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(57) Abstract: The present invention provides a method for detecting and differentiating disease states with high sensitivity and specificity. The method allows for a determination of whether a cellbased sample contains abnormal cells and, for certain diseases, is capable of determining the histologic type of disease present. The method detects changes in the level and pattern of expression of the molecular markers in the cell-based sample. Panel selection and validation procedures are also provided.

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CELL-BASED DETECTION AND DIFFERENTIATION OF DISEASE STATES

CROSS REFERENCE TO RELATED APPLICATIONS.

[0000.1] The present application is a continuation-in-part of U.S. Application Serial No. 10/095,298, filed March 12, 2002, which claims the benefit of U.S. Provisional Application Serial No. 60/274,638, filed March 12, 2001, the entire contents of which are incorporated by reference herein.

BACKGROUND OF THE INVENTION

[0001] The present invention relates to early detection of a general disease state in a patient. The present invention also relates to discrimination (differentiation) between specific disease states in their early and later stages.

[0002] Early detection of a specific disease state can greatly improve a patient's chance for survival by permitting early diagnosis and early treatment while the disease is still localized and its pathologic effects limited anatomically, physiologically, and clinically. Two key evaluative measures of any test or disease detection method are its sensitivity (Sensitivity = True Positives/(True Positives + False Negatives) and specificity (Specificity = True Negatives/(False Positives + True Negatives), which measure how well the test performs to accurately detect all affected individuals without exception, and without falsely including individuals who do not have the target disease. Historically, many diagnostic tests have been criticized due to poor sensitivity and specificity.

[0003] Sensitivity is a measure of a test's ability to detect correctly the target disease in an individual being tested. A test having poor sensitivity produces a high rate of false negatives, i.e., individuals who have the disease but are falsely identified as being free of that particular disease. The potential danger of a false negative is that the diseased individual will remain undiagnosed and untreated for some period of time, during which the disease may progress to a later stage wherein treatments, if any, may be less effective. This may result in poorer patient outcomes. An example of a test that has low sensitivity is a protein-based blood test for HIV. This type of test exhibits poor sensitivity because it fails to detect the presence of the virus until the disease is well established and the virus has invaded the bloodstream in substantial numbers.

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In contrast, an example of a test that has high sensitivity is viral-load detection using the polymerase chain reaction (PCR). High sensitivity is achieved because this type of test can detect very small quantities of the virus (see Lewis, D.R. et al. "Molecular Diagnostics: The Genomic Bridge Between Old and New Medicine: A White Paper on the Diagnostic Technology and Services Industry" Thomas Weisel Partners, June 13, 2001).

[0004] Specificity, on the other hand, is a measure of a test's ability to identify accurately patients who are free of the disease state. A test having poor specificity produces a high rate of false positives, i.e., individuals who are falsely identified as having the disease. A drawback of false positives is that they force patients to undergo unnecessary medical procedures treatments with their attendant risks, emotional and financial stresses, and which could have adverse effects on the patient's health. A feature of diseases which makes it difficult to develop diagnostic tests with high specificity is that disease mechanisms often involve a plurality of genes and proteins. Additionally, certain proteins may be elevated for reasons unrelated to a disease state. An example of a test that has high specificity is a gene-based test that can detect a p53 mutation. A p53 mutation will never be detected unless there are cancer cells present (see Lewis, D.R. et al. "Molecular Diagnostics: The Genomic Bridge Between Old and New Medicine: A White Paper on the Diagnostic Technology and Services Industry" Thomas Weisel Partners, June 13, 2001). [0005] Cellular markers are naturally occurring molecular structures within cells that can be discovered and used to characterize or differentiate cells in health and disease. Their presence can be detected by probes, invented and developed by human beings, which bind to markers enabling the markers to be detected through visualization and/or quantified using imaging systems. Four classes of cell-based marker detection technologies are cytopathology, cytometry, cytogenetics and proteomics, which are identified and described below.

[0006] Cytopathology relies upon the visual assessment by human experts of cytomorphological changes within stained whole-cell populations. An example is the cytological screening and cytodiagnosis of Papanicolaou-stained (i.e., Pap smear) cervical-vaginal specimens by cytotechnologists and cytopathologists, respectively. Unlike cytogenetics, proteomics and cytometry, cytopathology is not a quantitative tool. While it is the state-of-the-art in clinical diagnostic cytology, it is subjective and the diagnostic results are often not highly sensitive or reproducible, especially at early stages of cancer (e.g., ASCUS, LSIL).

[0007] Tests that rely on morphological analyses involve observing a sample of a patient's cells under an optical microscope to identify abnormalities in cell and nuclear shape, size, optical texture, or staining behavior. When viewed through a microscope, normal mature epithelial cells appear large and well differentiated, with condensed nuclei. Cells characterized by dysplasia, however, may be in a variety of stages of differentiation, with some cells being very immature. Finally, cells characterized by invasive carcinoma often appear undifferentiated, with very little cytoplasm and relatively large nuclei.

[0008] A drawback to diagnostic tests that rely on morphological analyses is that cell morphology is a lagging indicator. Since form follows function, often the disease state has already progressed to a critical, or advanced stage by the time the disease becomes evident by morphological analysis. The initial stages of a disease involve chemical changes at a molecular level. Changes that are detectable by viewing cell features under a microscope are typically not apparent until later stages of the disease. Therefore, tests that measure chemical changes on a molecular level, referred to as "molecular diagnostic" tests, are more likely to provide early detection than tests that rely on morphological analyses alone.

[0009] Cytometry is based upon the flow-microfluorometric instrumental analysis of fluorescently stained cells moving in single file in solution (flow cytometry) or the computer-aided microscope instrumental analysis of stained cells deposited onto glass microscope slides (image cytometry). Flow cytometry applications include leukemia and lymphoma immunophenotyping. Image cytometry applications include DNA ploidy, Malignancy-Associated Changes (MACs), cell-cycle kinetics and S-phase analyses. The flow and image cytometry approaches yield quantitative data characterizing the cells in suspension or on a glass microscope slide. Flow and image cytometry can produce good marker detection and differentiation results depending upon the sensitivity and specificity of the cellular stains and flow/image measurement features used.

[0010] Malignancy-Associated Changes (MACs) have been qualitatively observed and reported since the early to mid-1900's (OC Gruner: "Study of the changes met with leukocytes in certain cases of malignant disease" in Brit J Surg 3: 506-522, 1916) (HE Neiburgs, FG Zak, DC Allen, H Reisman, T Clardy: "Systemic cellular changes in material from human and animal tissues" in Transactions, 7th Ann Mtg Inter Soc Cytol Council, pp 137-144, 1959). From the mid-1900's through 1975, MACs were documented in independent qualitative histology and cytology studies

in buccal mucosa and buccal smears (Nieburgs, Finch, Klawe), duodenum (Nieburgs), liver (Elias, Nieburgs), megakaryocytes (Ramsdahl), cervix (Nieburgs, Howdon), skin (Kwitiken), blood and bone marrow (Nieburgs), monocytes and leukocytes (van Haas, Matison, Clausen), and lung and sputum (Martuzzi and Oppen Toth). Before 1975 these qualitative studies reported MAC-based sensitivities for specific disease detection from 76% to 97% and specificities from 50% to 90%. In 1975, Oppen Toth reported a sensitivity of 76% and specificity of 81% in a qualitative sputum analysis study.

[0011] Quantitative observations regarding MAC-based probe analysis began two to three decades ago (H Klawe, J Rowinski: "Malignancy associated changes (MAC) in cells of buccal smears detected by means of objective image analysis" in Acta Cytol 18: 30-33, 1974) (GL Wied, PH Bartels, M Bibbo, JJ Sychra: "Cytomorphometric markers for uterine cancer in intermediate cells" in Analyt Quant Cytol 2: 257-263, 1980) (G Burger, U Jutting, K Rodenacker: "Changes in benign population in cases of cervical cancer and its precursors" in Analyt Quant Cytol 3: 261-271, 1981). MACs were documented in independent quantitative histology and cytology studies in buccal mucosa and smears Klawe, Burger), cervix (Wied, Burger, Bartels, Vooijs, Reinhardt, Rosenthal, Boon, Katzke, Haroske, Zahniser), breast (King, Bibbo, Susnik), bladder and prostate (Sherman, Montironi), colon (Bibbo), lung and sputum (Swank, MacAulay, Payne), and nasal mucosa (Reith) studies with MAC-based sensitivities from 70% to 89% and specificities from 52% to 100%. Marek and Nakhosteen showed (1999, American Thoracic Society annual meeting) the results from two quantitative pulmonary (bronchial washings) studies showing (a) sensitivity of 89% and specificity of 92%, and (b) sensitivity of 91% and specificity of 100%.

[0012] Clearly, Malignancy-Associated Changes (MACs) are potentially useful probes that result from the image-cytometry marker detection technology. MAC-based features from DNA-stained nuclei can be used in conjunction with other molecular diagnostic probes to create optimized molecular diagnostic panels for the detection and differentiation of lung cancer and other disease states.

[0013] Cytogenetics detects specific chromosome-based intracellular changes using, for example, in situ hybridization (ISH) technology. ISH technology can be based upon fluorescence (FISH), multi-color fluorescence (M-FISH), or light-absorption-based chromogenics imaging (CHRISH) technologies. The family of ISH technologies uses DNA or

RNA probes to detect the presence of the complementary DNA sequence in cloned bacterial or cultured eukaryotic cells. FISH technology can, for example, be used for the detection of genetic abnormalities associated with certain cancers. Examples include probes for Trisomy 8 and HER-2 neu. Other highly sensitive as well as specific technologies such as polymerase chain reactions (PCR) can be used to detect B-cell and T-cell gene rearrangements. Cytogenetics is a highly specific marker detection technology since it detects the causative or "trigger" molecular event producing a pathology condition. It may, in general, be less sensitive than the other marker detection technologies because fewer events may be present to detect. In situ hybridization (ISH) is a molecular diagnostic method that uses gene-based analyses to detect abnormalities on the genetic level such as mutations, chromosome errors or genetic material inserted by a specific pathogen. For example, in situ hybridization may involve measuring the level of a specific mRNA by treating a sample of a patient's cells with labeled primers designed to hybridize to the specific mRNA, washing away unbound primers and measuring the signal of the label. Due to the uniqueness of gene sequences, a test involving the detection of gene sequences will likely have a high specificity, yielding very few false positives. However, because the amount of genetic material in a sample of cells may be very low, only a very weak signal may be obtained. Therefore, in situ hybridization tests that do not employ pre-amplification techniques will likely have a poor specificity, yielding many false negatives.

[0014] Proteomics depends upon cell characterization and differentiation resulting from the over-expression, under-expression, or presence/absence of unique or specific proteins in populations of normal or abnormal cell types. Proteomics includes not only the identification and quantification of proteins, but also the determination of their localization, modifications, interactions, chemical activities, and cellular/extracellular functions. Immunochemistry (IC) (immunocytochemistry in cells and immunohistochemistry (IHC) in tissues) is the technology used, either qualitatively or quantitatively (QIHC) to stain antigens (i.e., proteomes) using antibodies. Immunostaining procedures use a dye as the detection indicator. Examples of IHC applications include analyses for ER (estrogen receptor), PR progesterone receptor), p53 tumor suppressor genes, and EGRF prognostic markers. Proteomics is typically a more sensitive marker detection technology than cytogenetics because there are often orders of magnitude more protein molecules to detect using proteomics than there are cytogenetic mutations or genesequence alterations to detect using cytogenetics. However, proteomics may have a poorer

specificity than the cytogenetic marker detection technology since multiple pathologies may result in similar changes in protein over-expression or under-expression. Immunochemistry involves histological or cytological localization of immunoreactive substances in tissue sections or cell preparations, respectively, often utilizing labeled antibodies as probe reagents. Immunochemistry can be used to measure the concentration of a disease marker (specific protein) in a sample of cells by treating the cells with an agent such as a labeled antibody (probe) that is specific for an epitope on the disease marker, then washing away unbound antibodies and measuring the signal of the label. Immunochemistry is based on the property that cancer cells possess different levels of certain disease markers than do healthy cells. The concentration of a disease marker in a cancer cell is generally large enough to produce a large signal. Therefore, tests that rely on immunochemistry will likely have a high sensitivity, yielding few false negatives. However, because other factors in addition to the disease state may cause the concentration of a disease marker to become raised or lowered, tests that rely on immunochemical analysis of a specific disease marker will likely have poor specificity, yielding a high rate of false positives.

[0015] The present invention provides for a noninvasive disease state detection and discrimination method with both high sensitivity and high specificity. This method is useful for patient screening. The present invention also provides a disease state detection and discrimination method with both high sensitivity and high specificity. This method is useful for patient diagnosis and therapeutic monitoring. The method involves contacting a cytological sample or multiple samples suspected of containing diseased cells with a panel of probes comprising a plurality of agents, each of which quantitatively binds to a specific disease marker, and detecting and analyzing the pattern of binding of the probe agents. The present invention also provides methods of constructing and validating a panel of probes for detecting a specific disease (or group of diseases) and discriminating among its various disease states. Illustrative panels for detecting lung cancer and discriminating among different types of lung cancer are also provided. Illustrative panels or other cancers and non-cancer disease states are also provided. [0016] A human disease results from the failure of the human organism's adaptive mechanisms to neutralize external (i.e., local or global environmental) or internal insults which result in abnormal structures or functions within the body's cells, tissues, organs or systems. Diseases can be grouped by shared mechanisms of causation as illustrated below, in Table 1.

Table 1:

Classes of Diseases	Examples of Disease States
Allergy	Adverse reactions to foods and plants
Cardiovascular	Heart failure, atherosclerosis
Degenerative (neurological and muscular)	Alzheimer's and Parkinson's
Diet	Non-nutritional substances and excess/imbalanced nutrition
Hereditary	Sickle cell anemia, cystic fibrosis
Immune	HIV and autoimmune
Infection	Viral, bacterial, fungal, parasitic
Metabolic	Diabetes
Molecular and cell biology	Cancer (neoplasia)
Toxic insults	Alcohol, drugs, environmental mutagens and carcinogens
Trauma	Bodily injury from automobile collision

[0017] Disease states are either caused by or result in abnormal changes (i.e., pathological conditions) at a subcellular, cellular, tissue, organ, or human anatomic or physiological system level. Many disease states (e.g., lung cancer) are characterized by abnormal changes at a subcellular or cellular level. Specimens (e.g., cervical Pap smears, voided urine, blood, sputum, colonic washings) can be collected from patients with suspected disease states to diagnose those patients for the presence and type of the disease state. Molecular pathology is the discipline that attempts to identify and diagnostically exploit the molecular changes associated with these cell-based diseases.

[0018] Lung cancer is an illustrative example of a disease state in which screening of high-risk populations and at-risk individuals can be performed using diagnostic tests (e.g., molecular diagnostic panel assays) to detect the presence of the disease state. Also, for patients in which lung cancer or other disease states have been detected by these means, related diagnostic tests can

be employed to differentiate the specific disease state from related or co-occurring disease states. For example, in this lung cancer illustration, additional molecular diagnostic panel assays may indicate the probabilities that the patient's disease state is consistent with one of the following types of lung cancer: (a) squamous cell carcinoma of the lung, (b) adenocarcinoma of the lung, (c) large cell carcinoma of the lung, (d) small cell carcinoma of the lung, or (e) mesothelioma. Early detection and differentiation of cell-based disease states is a hypothesized means to improve patient outcomes.

[0019] Cancer is a neoplastic disease, the natural course of which is fatal. Cancer cells, unlike benign tumor cells, exhibit the properties of invasion and metastasis and are highly anaplastic. Cancer includes the three broad categories of carcinoma (i.e., epithelial cell-based cancers), sarcoma (e.g., bone-based cancers), and blood-based cancers (e.g., leukemia and lymphoma), but in lay usage each of the three types is often referred to synonymously with carcinoma. According to the World Health Organization (WHO), cancer affects more than 10 million people each year and is responsible for in excess of 6.2 million deaths.

[0020] Cancer is, in reality, a heterogeneous collection of diseases that can occur in virtually any part of the body. As a result, different treatments are not equally effective in all cancers or even among the stages of a specific type of cancer. Advances in diagnostics (e.g., mammography, cervical cytology, and serum PSA testing) have, in some cases, allowed for the detection of early-stage cancer when there are a greater number of treatment options, and therapies tend to be more effective. In cases where a solid tumor is small and localized, surgery alone may be sufficient to produce a cure. However, in cases where the tumor has spread, surgery may provide, at best, only limited benefits. In such cases the addition of chemotherapy and/or radiation therapy may be used to treat metastatic disease. While somewhat effective in prolonging life, treatment of patients with non-blood-based metastatic disease rarely produces a cure. Even through there may be an initial response, with time the disease progresses and the patient ultimately dies from its effects and/or from the toxic effects of the treatments.

[0021] While not proven, it is generally accepted that early detection and treatment will reduce

the morbidity, mortality and cost of cancer. Early detection will, in many cases, permit treatment to be initiated prior to metastasis. Furthermore, because there are a greater number of treatment options, there is a higher probability of achieving a cure or significant improvement in long-term survival.

[0022] Developing a test that can be used to screen an "at-risk" population has long been a goal of health practitioners. While there have been some successes such as mammography for breast cancer, PSA testing for prostate cancer, and the Pap smear for cervical cancer, in most cases cancer is detected at a relatively late stage where the patient is symptomatic and the disease is almost always fatal. For most cancers, no test or combination of tests has exhibited the necessary sensitivity and specificity to permit cost-effective identification of patients with early stage disease.

[0023] For a cancer screening program to be successful and gain acceptance by patients, physicians, and third-party payers, the test must have implied benefit (changes the outcome), be widely available and be able to be carried out readily within the framework of general healthcare. The test should be relatively noninvasive, leading to adequate compliance, have high sensitivity, and reasonable specificity and predictive value. In addition, the test must be available at relatively low cost.

[0024] For patients who are suspected of having cancer, the diagnosis must be confirmed and the tumor properly staged cytologically and clinically in order for physicians to undertake appropriate therapeutic intervention. Some tests currently being used in the diagnosis and staging of cancer, however, either lack sufficient sensitivity or specificity, are too invasive, or are too costly to justify their use as a population-based screening test. Shown below in Tables 2 and 3, for example, are estimates of sensitivity and specificity of lung cancer diagnostics and estimated costs (U.S. dollars) for diagnostic tests used to detect lung cancer.

Table 2:

ESTIMATES OF SENSITIVITY AND SPECIFICITY OF LUNG CANCER DIAGNOSTICS [1]

DIACNOSTIC TEST	SENSTITVITY (%)	SPECIFICITY (%).
Conventional Sputum Cytology	51.0	100.0
Chest X-ray	16-85*	90-95
White Light Bronchoscopy	48.0-80.0	91.1-96.8
LIFE Bronchoscopy	72.0	86.7
Computed Tomography	63.0-99.9	80.0-61
PET Scan	88.0-92.5	83.0-93.0

^{*}Dependent upon the stage of the disease at the time of diagnosis

Table 3:

ESTIMATED COSTS FOR DIAGNOSTIC TESTS USED IN LUNG CANCER [1]

DIAGNOSTIC TEST	COST (\$)
Sputum Cytology	90
Chest X-ray	. 44
Bronchoscopy	725
Computed Tomography	378
PET Scan	800-3000
Open Biopsy	12,847-14,121

[0025] The chest radiograph (X-ray) is often used to detect and localize cancer lesions due to its reasonable sensitivity, high specificity and low cost. However, small lesions are often difficult to detect and although larger tumors are relatively easy to visualize on a chest film, at the time of detection most have already metastasized. Thus, chest X-rays lack the necessary sensitivity for use as an early detection method.

[0026] Computed tomography (CT) is useful in the confirmation and characterization of pulmonary nodules and allows the detection of subtle abnormalities that are often missed on a standard chest X-ray [2]. CT, and Spiral CT methods in particular, remains the test of choice for patients who present with a prior malignant sputum cytology result or vocal chord paralysis. CT, with its improved sensitivity over the conventional chest film, has become the primary tool for imaging the central airway [3]. While capable of examining large areas, CT is subject to artifacts from cardiac and respiratory motion although improved resolution can be achieved through the use of iodinated contrast material.

[0027] Spiral CT is a more rapid and sensitive form of CT that has the potential to detect early cancer lesions more reliably than either conventional CT or X-ray. Spiral CT appears to have greatly improved sensitivity in diagnosing early disease. However, the test has relatively low specificity with a 20% false positive rate [4]. As the resolution of Spiral CT instruments improve by engineering technology advances, the false positive rate is likely to increase. Spiral CT is also less sensitive in detecting the central lesions that represent one-third of all lung cancers. Furthermore, while the cost of the initial test is relatively low (\$300), the cost of follow-up can be at least an order of magnitude higher. Cytology using molecular diagnostic panel assays offers significant promise as an adjunctive test with Spiral CT to improve the specificity of Spiral

CT testing by minimizing false positive results through the evaluation of fine needle aspirations (FNAs) or biopsies (FNBs) from Spiral CT-suspicious pulmonary nodules.

[0028] Fluorescence bronchoscopy provides increased sensitivity over conventional white light bronchoscopy, significantly improving the detection of small lesions within the central airway [5]. However, fluorescence bronchoscopy is unable to detect peripheral lesions, it takes a long time for bronchoscopists to examine a patient's airways, and it is an expensive procedure. Additionally, the procedure is moderately invasive, creating an insurmountable barrier to its use as a population-based screening test.

[0029] Positron Emission Tomography (PET) is a highly sensitive test that utilizes radioactive glucose to identify the presence of cancer cells within the lung [6-8]: The cost of establishing a testing facility is high and there is the need for a cyclotron on site or nearby. Also, implementing centralized testing is a logistical problem. This, coupled with the high cost of the test, has limited the use of PET scans to staging lung cancer patients rather than for early detection of the disease. [0030] Although used for some time as a means of screening for lung cancer, sputum cytology has enjoyed only limited success due to its low sensitivity and its failure to reduce disease-specific mortality. In conventional sputum cytology, the pathologist uses characteristic changes in cellular morphology to identify malignant cells and make a diagnosis of cancer. Today only 15% of patients who are 'at-risk" or who are suspected of having lung cancer undergo sputum cytology testing, and less than 5% undergo multiple evaluations [9]. A number of factors including tumor size, location, degree of differentiation, cell clumping, inefficiency of clearing mechanisms to release cells and sputum to the external environment, and the poor stability of cells within the sputum contribute to the overall poor performance of the test.

[0031] Cancer diagnostics has traditionally relied upon the detection of single molecular markers. Unfortunately, cancer is a disease state in which single markers have typically failed to detect or differentiate many forms of the disease. Thus, probes that recognize only a single marker have been shown to be largely ineffective. Exhaustive searches for "magic bullet" diagnostic tests have been underway for many decades though no universal successful magic bullet probes have been found to date.

[0032] A major premise of this invention is that cell-based cancer diagnostics and the screening, diagnosis for, and therapeutic monitoring of other disease states will be significantly improved over the state-of-the-art that uses single marker/probe analyses rather than kits of

multiple, simultaneously labeled probes. This multiplexed analytical approach is particularly well suited for cancer diagnostics since cancer is not a single disease. Furthermore, this multifactorial "panel" approach is consistent with the heterogeneous nature of cancer, both cytologically and clinically.

[0033] Key to the successful implementation of a panel approach to cell-based diagnostic tests is the design and development of optimized panels of probes that can chemically recognize the pattern of markers that characterizes and distinguishes a variety of disease states. This patent application describes an efficient and unique methodology to design and develop such novel and optimized panels.

[0034] Improved methods for specimen collection (e.g., point-of-care mixers for sputum cytology) and preparation (e.g., new cytology preservation and transportation fluids, and liquid-based cytology preparation instruments) are under development and becoming commercially available. In conjunction with existing and these emerging methods, a successful implementation of this molecular diagnostics cell-based panel assay will lead to (a) characterization of the molecular profile of malignant tumors and other disease states, (b) improved methods for early cancer and other disease state detection and differentiation, and (c) opportunities for improved clinical diagnoses, prognoses, customized patient treatments, and therapeutic monitoring.

SUMMARY OF THE INVENTION

[0035] The present invention is directed to a panel for detecting a generic disease state or discriminating between specific disease states using cell-based diagnosis. The panel comprises a plurality of probes each of which specifically binds to a marker associated with a generic or specific disease state, wherein the pattern of binding of the component probes of the panel to cells in a cytology specimen is diagnostic of the presence or specific nature of said disease state. The present invention is also directed to a method of forming a panel for detecting a disease state or discriminating between disease states in a patient using cell-based diagnosis. The method involves determining the sensitivity and specificity of binding of probes each of which specifically binds to a member of a library of markers associated with a disease state and selecting a limited plurality of said probes whose pattern of binding is diagnostic for the presence or specific nature of said disease state. The present method is also directed to a method of detecting a disease or discriminating between disease states. The method involves contacting a cytological sample suspected of containing abnormal cells characteristic of a disease state with a

panel according to claim 1 and detecting a pattern of binding of said probes that is diagnostic for the presence or specific nature of said disease state.

BRIEF DESCRIPTION OF THE FIGURES

[0036] Figure 1. Molecular markers that are preferable markers to be included in a panel for identifying different histologic types of lung cancer. The column labeled "%" indicates the percentage of tumor specimens that express a particular marker.

Potential ways in which different markers may be used to discriminate [0037] Figure 2. between specific types of lung cancer. SQ indicates squamous cell carcinoma, AD indicates adenocarcinoma, LC indicates large cell carcinoma, SC indicates small cell carcinoma and ME indicates mesothelioma. The numbers appearing in each cell represent frequency of marker change in one cell type versus another. To be included in the table, the ratio must be greater than 2.0 or less than 0.5. A number larger than 100 generally indicates that the second marker is not expressed. In such cases the denominator was set at 0.1 for the purpose of the analysis. Finally, empty cells represent either no difference in expression or the absence of expression data. Comparisons between H-scores for probes 7 and 15 in control tissue and in [0038] Figure 3. cancerous tissue. The X-axis shows the H-scores while the Y-axis shows the percent of cases. Correlation matrix, in which correlation measures the amount of linear [0039] Figure 4. association between a pair of variables. All markers in this matrix with a correlation number of 50% or higher are considered correlate markers. Note that all diagonal elements of this correlation matrix have a value of 1.0 (i.e., True) because the diagonal elements show autocorrelation values (i.e., Probe N correlation to Probe N). Also, note that this matrix is diagonally symmetric (i.e., correlation value of Probe N versus M is identical to the correlation value of Probe M versus N).

[0040] Figure 5. Detection panel compositions, pair-wise discrimination panel compositions and joint discrimination panel compositions. Panel compositions using decision tree analysis, stepwise LR and stepwise LD are shown. Note that shaded boxes identify probes that are shown to be effective by two or more of these independent analytical methods.

[0041] Figure 6. Detection panel compositions wherein probe 7 was not included as a probe. Panel compositions using decision tree analysis, stepwise LR and stepwise LD are shown. Note that shaded boxes identify probes that are shown to be effective by two or more of these independent analytical methods.

[0042] Figure 7. Detection panel compositions using only commercially preferred probes. Panel compositions using decision tree analysis, stepwise LR and stepwise LD are shown. Note that shaded boxes identify probes that are shown to be effective by two or more of these independent analytical methods.

[0043] Figures 8a-c. Summary of the preferred markers (probes) for panels for detecting and/or diagnosing lung, colorectal, bladder, prostate, breast and cervical cancer.

DETAILED DESCRIPTION OF THE INVENTION

1. Introduction

[0044] The present invention provides a noninvasive disease state detection and discrimination method with high sensitivity and specificity. The method involves contacting a cytological or histological sample or sample suspected of containing diseased cells with a panel comprising a plurality of agents, each of which quantitatively binds to a disease marker, and detecting a pattern of binding of the agents. This pattern includes the localization and density/concentration of binding of the component probes of the panel. The present invention also provides methods of making a panel for detecting a disease and also for discriminating between disease states as well as panels for detecting lung cancer in early stages and discriminating between different types of lung cancer. Panel tests have been used in medicine. For example, panels are used in blood serum analysis. However, because a cytology analysis involves imaging and localization of specific markers within individual cells and tissues, prior to the present invention it was not apparent that the panel approach would be effective for cytology or histology samples. Additionally, it was not apparent which, if any statistical analyses could be applied to design and develop an optimized cell-based diagnostic panel of probes.

[0045] One of the few examples of a cytology-based screening program is the Pap Smear, which screens for cervical cancer. For over 50 years this method has been practiced and has greatly contributed to the fact that today, almost no woman who has regular Pap smears dies of cervical cancer. There are drawbacks, however, to the Pap smear screening program. For example, Pap smears are labor intensive, subject to the variability associated with human performance, and are not universally accessible. The present molecular diagnostic cell-based screening method utilizing probe panels does not suffer from these drawbacks. The method may be fully automated and thereby made less expensive and reproducible, increasing access to this type of testing.

[0046] The present invention provides a method, having both high specificity and high sensitivity, for detecting a disease state and for discriminating between disease states. The invention is applicable to any cell-based disease state, such as cancer and infectious diseases.

[0047] The panel is diagnostic of the presence or specific nature of the disease state. The present invention overcomes the limitations and drawbacks of known disease state detection methods by enabling quick, accurate, relatively noninvasive and easy detection and

[0048] A feature of the inventive method for making a panel of the present invention is the rapidity with which the panel may be developed.

discrimination of diseased cells in a cytological sample while keeping costs low.

[0049] There are several benefits to using a panel of agents in a method for detecting a disease state, and for discriminating between types of disease states. One benefit is that a panel of agents has sufficient redundancy to permit detection and characterization of disease states thereby increasing the sensitivity and specificity of the test. Given the heterogeneous nature of many disease states, no single agent is capable of identifying the vast majority of cases.

[0050] An additional benefit to using a panel is that use of a panel permits discrimination between the various types of a disease state based on specific patterns (probe localization and density/concentration) of expression. As the various types of a disease may exhibit dramatic differences in their rate of progression, response to therapy, and lethality, knowledge of the specific type can help physicians choose the optimal therapeutic approach.

2. The Panel

[0051] The panel of the present invention comprises a plurality of agents, each of which quantitatively binds to a disease marker, wherein the pattern (localization and density/concentration) of binding of the component agents of the panel is diagnostic of the presence or specific nature of a disease state. Therefore, the panel may be a detection panel or a discrimination panel. A detection panel detects whether a generic disease state is present in a sample of cells, while a discrimination panel discriminates among different specific disease states in a sample of cells known to be affected by a disease state which comprises different types of diseases. The difference between a detection panel and a discrimination panel lies in the specific agents that the panels comprise. A detection panel comprises agents having a pattern of binding that is diagnostic of the presence of a disease state, while a discrimination panel comprises agents

having a pattern of binding that allows for determining the specific nature (i.e., each type) of the disease state.

[0052] A panel, by definition, contains more than one member. There are several reasons why it is beneficial to use a panel of markers rather than just one marker alone to detect a generic disease state or to discriminate among specific disease states. One reason is the unlikely existence of a probe for one single marker, that is present in all diseased cells yet not present in healthy cells, whose behavior can be measured with a high specificity and sensitivity to yield an accurate test result. If such a single probe existed for detection of a particular disease with high sensitivity and specificity, it would already have been utilized for clinical testing. Rather, it is the directed selection of panel tests, each consisting of multiple probes, that together can provide the range of detection capability to ensure clinically adequate testing.

[0053] If one nevertheless chooses to construct a panel test comprising one or a very few probes, then the failure of any single marker/probe combination to perform its labeling function for any reason (for example, diminished reactivity of the specimen cells due to biological variability; inherent variability between lots of probe reagents; a weak, outdated or defective processing reagent; improper processing time or conditions for that probe) could result in a catastrophic failure of the test to detect or discriminate the target disease. The inclusion of multiple, and even redundant probes in each panel test greatly enhances the probability that a failure of any one probe will not cause a catastrophic failure of the test.

[0054] A probe is any molecular structure or substructure that binds to a disease marker. The term "agent" as used herein, may also refer to a molecular structure or substructure that binds to a disease marker. Molecular probes are homing devices used by biologists and clinicians to detect and locate markers indicative of the specific disease states. For example, antibodies may be produced that bind specifically to a protein previously identified as a marker for small cell lung cancer. This antibody probe can then be used to localize the target protein marker in cells and tissues of patients suspected of having the disease by using appropriate immunochemical protocols and incubations. If the antibody probe binds to its target marker in a stoichiometric (i.e., quantitative) fashion and is labeled with a chromogenic or colored "tag", then localization and quantitation of the probe and, indirectly, its target marker may be accomplished using an optical microscope and image cytometry technology.

[0055] The present invention contemplates detecting changes in molecular marker expression at the DNA, RNA or protein level using any of a number of methods available to an ordinary skilled artisan. Exemplary probes may be a polyclonal or monoclonal antibody or fragment thereof or a nucleic acid sequences that is complementary to the nucleic acid sequence encoding a molecular marker in the panel. A probe may also be a stain, such as a DNA stain. Many of the antibodies used in the present invention are specific to a variety of cell surface or intracellular antigens as marker substances. The antibodies may be synthesized using techniques generally known to those of skill in the art. For example, after the initial raising of antibodies to the marker, the antibodies can be sequenced and subsequently prepared by recombinant techniques. Alternatively, antibodies may be purchased.

[0056] In embodiments of the present invention, the probe contains a label. A probe containing a label is often referred to herein as a "labeled probe". The label may be any substance that can be attached to a probe so that when the probe binds to the marker a signal is emitted or the labeled probe can be detected by a human observer or an analytical instrument. This label may also be referred to as a "tag". The label may be visualized using reader instrumentation. The term "reader instrumentation" refers to the analytical equipment used to detect a probe. Labels envisioned by the present invention are any labels that emit a signal and allow for identification of a component in a sample. Preferred labels include radioactive, fluorogenic, chromogenic or enzymatic moieties. Therefore, possible methods of detection include, but are not limited to, immunocytochemistry, immunohistochemistry, in situ hybridization, fluorescent in situ hybridization, flow cytometry and image cytometry. The signal generated by the labeled probe is of sufficient intensity to permit detection by a medical practitioner.

[0057] A "marker", "disease marker" or "molecular marker" is any molecular structure or substructure that is correlated with a disease state or pathogen. The term "antigen" may be used interchangeably with "marker". Broadly defined, a marker is a biological indicator that may be deliberately used by an observer or instrument to reveal, detect, or measure the presence or frequency and/or amount of a specific condition, event or substance. For example, a specific and unique sequence of nucleotide bases may be used as a genetic marker to track patterns of genetic inheritance among individuals and through families. Similarly, molecular markers are specific molecules, such as proteins or protein fragments, whose presence within a cell or tissue indicates a particular disease state. For example, proliferating cancer cells may express novel cell-surface

proteins not found on normal cells of the same type, or may over-express specific secretory proteins whose increased or decreased abundance (e.g., overexpression or underexpression, respectively) can serve as markers for a particular disease state.

[0058] Suitable markers for cytology panels are substances that are localized in or on the nucleus, cytoplasm or cell membrane. Markers may also be localized in organelles located in any of these locations in the cell. Exemplary markers localized in the nucleus include but are not limited to retinoblastoma gene product (Rb), Cyclin A, nucleoside diphosphate kinase/nm23, telomerase, Ki-67, Cyclin D1, proliferating cell nuclear antigen (PCNA), p120 (proliferationassociated nucleolar antigen) and thyroid transcription factor 1 (TTF-1). Exemplary markers localized in the cytoplasm include but are not limited to VEGF, surfactant apoprotein A (SP-A), nucleoside nm23, melanoma antigen-1 (MAGE-1), Mucin 1, surfactant apoprotein B (SP-B), ER related protein p29 and melanoma antigen-3 (MAGE-3). Exemplary markers localized in the cell membrane include but are not limited to VEGF, thrombomodulin, CD44v6, E-Cadherin, Mucin 1, human epithelial related antigen (HERA), fibroblast growth factor (FGF), heptocyte growth factor receptor (C-MET), BCL-2, N-Cadherin, epidermal growth factor receptor (EGFR) and glucose transporter-3 (GLUT-3). An example of a marker located in an organelle of the cytoplasm is BCL-2, located (in part) in the mitochondrial membrane. An example of a marker located in an organelle of the nucleus is p120 (proliferating-associated nucleolar antigen), located in the nucleoli.

[0059] Preferred are markers where changes in expression: occur early in disease progression, are exhibited by a majority of diseased cells, allow for detection of in excess of 75% of a given disease type; most preferably in excess of 90% of a given disease type and/or allow for the discrimination between the nature of different types of a disease state.

[0060] It is noted that the inventive panel may be referred to as a panel of probes or a panel of markers, since the probes bind to the markers. Therefore, the panel may comprise a number of markers or it may comprise a number of probes that bind to specific markers. For the sake of consistency, the present panel is referred to as a panel of probes; however, it could also be referred to as a panel of markers.

[0061] Markers can also include features such as malignancy-associated changes (MACs) in the cell nucleus or features related to the patient's family history of cancer. Malignancy-associated changes, or MACs, are typically sub-visual changes that occur in normal-appearing

cells located in the vicinity of cancer cells. These exceedingly subtle changes in the cell nucleus may result biologically from changes in the nuclear matrix and the chromatin distribution pattern. They cannot be appreciated even by trained observers through the visual observation of individual cells, but may be determined from statistical analysis of cell populations using highly automated, computerized high-speed image cytometry. Techniques for detection of MACs are well known to those of skill in the art and are described in more detail in: Gruner, O.C. Brit J. Surg. 3 506-522 (1916); Neiburgs, H.E. et al., Transaction, 7th Annual Mtg. Inter. Soc. Cytol. Council 137-144 (1959); Klawe, H. Acta. Cytol. 18 30-33 (1974); Wied, G.L., et al., Analty. Quant. Cytol. 2 257-263 (1980); and Burger, G., et al., Analyt. Quant. Cytol. 3 261-271 (1981). The present invention encompasses any marker that is correlated with a disease state. The individual markers themselves are mere tools of the present invention. Therefore, the invention is not limited to specific markers. One way to classify markers is by their functional relationship to other molecules. As used herein, a "functionally related" marker is a component of the same biological process or pathway as the marker in question and would be known by a person of skill in the art to be abnormally expressed together with the marker in question. For example, many markers are associated with a cell proliferation pathway, such as fibroblast growth factor (FGF), (vascular endothelial growth factor) VEGF, CyclinA and Cyclin D1. Other markers are glucose transporters, such as Glut-1 and Glut-3.

[0063] A person of ordinary skill in the art is well equipped to determine a functionally related marker and may research various markers or perform experiments in which the functional behavior of a marker is determined. By way of non-limiting example, a marker may be classified as a molecule involved in angiogenesis, a transmembrane glycoprotein, a cell surface glycoprotein, a pulmonary surfactant protein, a nuclear DNA-binding phosphoprotein, a transmembrane Ca²⁺ dependent cell adhesion molecule, a regulatory subunit of the cyclin-dependent kinases (CDK's), a nucleoside diphosphate kinase, a ribonucleoprotein enzyme, a nuclear protein that is expressed in proliferating normal and neoplastic cells, a cofactor for DNA polymerase delta, a gene that is silent in normal tissues yet when it is expressed in malignant neoplasms is recognized by autologous, tumor-directed and specific cytotoxic T cells (CTL's), a glycosylated secretory protein, the gastrointestinal tract or genitourinary tract, a hydrophobic protein of a pulmonary surfactant, a transmembrane glycoprotein, a molecule involved in proliferation, differentiation and angiogenesis, a proto-oncogene, a homeodomain transcription

factor, a mitochondrial membrane protein, a molecule found in nucleoli of a rapidly proliferating cell, a glucose transporter, or an estrogen-related heat shock protein.

[0064] Classes of biomarkers and probes include, but are not limited to: (a) morphologic biomarkers, including DNA ploidy, MACs and premalignant lesions; (b) genetic biomarkers including DNA adducts, DNA mutations and apoptotic indices; (c) cell cycle biomarkers including cellular proliferation, differentiation, regulatory molecules and apoptosis markers, and; (d) molecular and biochemical biomarkers including oncogenes, tumor suppressor genes, tumor antigens, growth factors and receptors, enzymes, proteins, prostaglandin levels and adhesion molecules.

[0065] A "disease state" may be any cell-based disease. In some embodiments the disease state is cancer. In other embodiments, the disease state is an infectious disease. The cancer may be any cancer, including, but not limited to epithelial cell-based cancers from the pulmonary, urinary, gastrointestinal, and genital tracts; solid and/or secretory tumor-based cancers, such as sarcomas, breast cancer, cancer of the pancreas, cancer of the liver, cancer of the kidneys, cancer of the thyroid, and cancer of the prostate; and blood-based cancers, such as leukemias and lymphomas. Exemplary cancers which may be detected by the present invention are lung, bladder, gastrointestinal, cervical, breast or prostate cancer. Exemplary infectious diseases which may be detected are cell-based diseases in which the infectious organism is a virus, bacteria, protozoan, parasite, or fungus. The infectious disease, for example, may be HIV, hepatitis, influenza, meningitis, mononucleosis, tuberculosis and sexually transmitted diseases (STDs), such as chlamydia, trichomonas, gonorrhea, herpes and syphilis.

[0066] As used herein, the term "generic disease state" refers to a disease which comprises several types of specific diseases, such as lung cancer, sexually transmitted diseases and immune-based diseases. Specific disease states are also referred to as histologic types of diseases. For example, the term "lung cancer" comprises several specific diseases, among which are squamous cell carcinoma, adenocarcinoma, large cell carcinoma, small cell lung cancer and mesothelioma. The term "sexually transmitted diseases" comprises several specific diseases, among which are Gonorrhea, Human Papilloma Virus (HPV), herpes and Syphilis. The term "immune-based diseases" comprises several specific diseases, such as systemic lupus erythematosus (Lupus), rheumatoid arthritis and pernicious anemia.

[0067] As used herein, the term "high-risk population" refers to a group of individuals who are exposed to disease causing agents, e.g., carcinogens, either at home or in the workplace (i.e., a "high risk population" for lung cancer might be exposed to smoking, passive smoking and occupational exposure). Individuals in a "high-risk population" may also have a genetic predisposition.

[0068] The term "at-risk" refers to individuals who are asymptotic but, because of a family history or significant exposure are at a significant risk of developing a disease state (i.e., an individual at risk for lung cancer with a > 30 pack-year history of smoking; "pack-year" is a measurement unit computed by multiplying the number of packs smoked per day, times the number of years for this exposure).

[0069] Cancer is a disease in which cells divide without control due to, for example, altered gene expression. In the methods and panels of the present invention, the cancer may be any malignant growth in any organ. For example, the cancer may be lung, bladder, gastrointestinal, cervical, breast or prostate cancer. Each cancer may comprise a collection of diseases or histological types of cancer. The term "histologic type" refers to cancers of different histology. Depending on the cancer there can be one or several histologic types. For example, lung cancer includes, but is not limited to, squamous cell carcinoma, adenocarcinoma, large cell carcinoma, small cell carcinoma and mesothelioma. Knowledge of the histologic type of cancer affecting a patient is very useful because it helps the medical practitioner to localize and characterize the disease and to determine the optimal treatment strategy.

[0070] Infectious diseases include cell-based diseases in which the infectious organism is a virus, bacteria, protozoan, parasite or fungus.

[0071] Exemplary detection and discrimination panels are panels that detect lung cancer, a general disease state, and panels that discriminate a single lung cancer type, specific disease state, against all other types of lung cancer and false positives. False positives can include metastatic cancer of a different type, such as metastasized liver, kidney or pancreatic cancer.

3. Methods of Making a Panel

[0072] The method of making a panel for detecting a generic disease state or discriminating between specific disease states in a patient involves determining the sensitivity and specificity of binding of probes to a library of markers associated with a generic or specific disease state and selecting a plurality of said probes whose pattern of binding (localization and

density/concentration) is diagnostic of the presence or specific nature of the disease state. In some embodiments, optional preliminary pruning and preparation steps are performed. The method of making a panel of the present invention involves analyzing the pattern of binding of probes to markers in known histologic pathology samples, i.e. gold standards. The classifier designed on the gold standard data can then be used to design a classifier for cytometry, especially automated cytometry. Therefore, the set of marker probes selected from the pathology analysis is used to prepare a new training data set taken from a cytology sample, such as sputum, fine needle aspirations, urine, etc. Cells shed from the specified lesions will stain in a similar fashion to the gold standards. The method described here eliminates the experimental error in selecting the best features set because the integrity of the diagnosis based on gold standard histologic pathology samples is high. Although it is, in principle, possible to use cytology samples to produce a panel, this is less desirable because cytology samples contain debris, there may be deterioration of the cells in a cytology sample, and the pathology diagnosis may be difficult to confirm clinically.

[0073] A library of markers is a group of markers. The library can comprise any number of markers. However, in some embodiments the number of markers in the library is limited by technical and/or commercial practicalities, such as specimen size. For example, in some embodiments, each specimen is tested against all of the markers in the panel. Therefore, the number of markers must not be larger than the number of samples into which the specimen may be divided. Another technical practicality is time. Typically, the library contains less than 60 markers. Preferably, the library contains less than 50 markers. More preferably, the library contains less than 40 markers. Most preferably the library contains 10-30 markers. It is preferable that the library of potential panel members contain more than 10 markers so that there is opportunity to optimize the performance of the panel. As used herein, the term "about" means plus or minus 3 markers.

[0074] In some embodiments, a library is obtained by consulting sources which contain information about various markers and correlations between the markers and generic/specific disease states. Exemplary sources include experimental results, theoretical or predicted analyses and literary sources, such as journals, books, catalogues and web sites. These various sources may use histology or cytology and may rely on cytogenetics, such as *in situ* hybridization; proteomics, such as immunohistochemistry; cytometry, such as MACs or DNA ploidy; and/or

cytopathology, such as morphology. The markers may be localized anywhere in or on a cell. For example, the markers may be localized in or on the nucleus, the cytoplasm or the cell membrane. The marker may also be localized in an organelle within any of the aforementioned localizations. [0075] In some embodiments, the library may be of an unsuitable size. Therefore, one or more pruning steps may be required prior to initiating the basic method for making a panel. The pruning step may involve one or several successive pruning steps. One pruning step may involve, for example, setting an arbitrary threshold for sensitivity and/or specificity. Therefore, any marker whose experimental or predicted sensitivity and/or specificity falls below the threshold may be removed from the library. Other exemplary pruning steps, which may be performed alone or in sequence with other pruning steps, may rely on detection technology requirements, access constraints and irreproducibility of reported results. With respect to detection technology requirements, it is possible that the machinery required to detect a particular marker is unavailable. With respect to access constraints, it is possible that licensing restrictions make it difficult or impossible to obtain a probe that binds to a particular marker. In some embodiments, a due diligence study is performed on each marker.

loopoil In some embodiments, prior to beginning the basic method for making a panel, it may be necessary to perform preparation steps. Exemplary preparation steps include optimizing the protocols for objective quantitative detection of the markers in the library and collecting histology specimens. Optimization of the protocols for objective quantitative detection of the markers is within the skill of an ordinary artisan. For example, the necessary reagents and supplies must be obtained, such as buffers, reagents, software and equipment. It is possible that the concentration of reagents may need to be adjusted. For example, if non-specific binding is observed, a person of ordinary skill in the art may dilute the concentration of the probe solution. [0077] In some embodiments, the histology specimens are Gold Standards. The term "Gold Standard" is known by a person of ordinary skill in the art to mean that the histology and clinical diagnosis of the specimen is known. The gold standards are often referred to as a "training" data set. The gold standards comprise a set of measurements, or reliable estimates, of all the features that may contribute to the discriminating process. Such features are collected from samples collected from a representative number of patients with known disease states. The standard samples can be cytology samples but this is less desirable for panel selection.

[0078] The histology samples may be obtained by any technique known to those of skill in the art, for example biopsy. In some embodiments, it is necessary that the size of the specimen per patient be large enough so that enough tissue sections can be obtained to test each marker in the library.

[0079] In some embodiments, specimens are obtained from multiple patients diagnosed with each specific disease state. One specimen per patient may be obtained, or multiple specimens per patient may be obtained. In embodiments in which multiple specimens are obtained from individual patients, the expertise of the surgeon is relied upon to establish that each specimen obtained from a single patient is similar to the other specimens obtained from that patient. Specimens are also obtained from a control group of patients. The control group of patients may be healthy patients or patients that are not suffering from the generic or specific disease state that is being tested.

[0080] The first step of the basic method is determining the sensitivity and specificity of binding of probes to a library of markers associated with the desired disease state. In this step, a probe that is specific for each marker in the library is applied to a sample of the patients' specimens. Therefore, in some embodiments, if there are, for example, 30 markers in the library, each patient's specimen will be divided into 30 samples and each sample will be treated with a probe that is specific for one of the 30 markers. The probe contains a label that may be visualized. Therefore, the pattern and level of binding of the probe to the marker can be detected. The pattern and level of binding may be detected either quantitatively, i.e., by an analytical instrument, or qualitatively, by a human, such as a pathologist.

[0081] In some embodiments, an objective and/or quantitative scoring method is developed to detect the pattern and level of binding of the probe to the markers. The scoring method may be heuristically designed. Scoring methods are used to objectify a subjective interpretation, for example, by a pathologist. It is within the skill of an ordinary artisan to determine a suitable scoring method. In some embodiments, the scoring method may comprise categorizing features, such as the density of a marker probe stain as: none, weak, moderate, or intense. In another embodiment, these features may be measured with algorithms operating on microscope slide images. An exemplary scoring method is one in which the proportions and density are consolidated into a single "H Score" obtained by grading the intensity as: none = 0, weak = 1, moderate = 2, intense = 3, and the percentage cells as: 0-5% = 0, 6-25% = 1, 26-50% = 2, 51-

75% = 3, >75% = 4, and then multiplying the two grades together. For example, 50% weakly stained plus 50% moderately stained would score $6 = (1 \times 2) + (2 \times 2)$. The "H score" honors the late Kenneth Hirsch, one of the present inventors.

[0082] An ordinary artisan is capable of addressing issues related to minimizing potential biases related to pathologists and samples. For example, randomizing may be used to minimize the chance of having a systematic error. Blinding may be used to eliminate experimental biases by the people conducting the experiments. For example, in some embodiments, pathologist-to-pathologist variation may be minimized by conducting a double blind study. As used herein, the term "double blind study" is a well establish method for avoiding biases, where the data collection and data analysis are done independently. In other embodiments, sample-to-sample variation is minimized by randomizing the samples. For example, the samples are randomized before the pathologist analyzes them. There is also randomization involved in the experimental protocols. In some embodiments, each sample is analyzed by at least two pathologists. For each patient, a reliable assessment of the binding of the probe to the marker is obtained. In one embodiment, this diagnosis is made by qualified pathologists, using two pathologists per patient, to check for reliability.

[0083] A sufficient number of samples should be collected to produce reliable designs and reliable statistical performance estimates. It is within the skill of a normal artisan to determine how many samples are sufficient to produce reliable designs and reliable statistical performance estimates. Most standard classifier design packages have methods for determining the reliability of the performance estimates and the sample size should be progressively increased until reliable estimates are achieved. For example, sufficient estimates to produce reliable designs may be achieved with 200 samples collected and 27 different features estimated from each sample. [0084] The second step is selecting a limited plurality of probes. The selecting step may employ statistical analysis and/or pattern recognition techniques. In order to perform the selecting step, the data may be consolidated into a database. In some embodiments, the probes may be numbered to render their method of action as unseen during the analysis of their effectiveness and further minimize biases. Rigorous statistical techniques are used because of the large amount of data that is generated by this method. Any statistical method may be used and an ordinary skilled statistician will be able to identify which and how many methods are appropriate.

[0085] Any number of statistical analysis and/or pattern recognition methods may be employed. Since the structure of the data is initially unknown, and since different classifier design methods perform better for different structures, it is preferred to use at least two design methods on the data. In some embodiments, three different methodologies may be used. One of ordinary skill in the art of statistical analysis and/or pattern recognition of data sets would recognize from characteristics of the data set structures that certain statistical methods would be more likely to yield an efficient result than others, where efficient in this case means achieving a certain level of sensitivity and specificity with a desired number of probes. A person of ordinary skill in the art would know that the efficiency of the statistical analysis and/or method is data dependent. Exemplary statistical analysis and/or pattern recognition methods are described below:

- [0086] a) A Decision Tree Method, known as C4.5. C4.5 is public domain software available via ftp from http://www.cse.unsw.edu.au/~quinlan/. This is well suited to data that can be best classified by sequentially applying a decision threshold to specific features in turn. This works best with uncorrelated data; it also copes with data with similar means provided the variances differ. The C4.5 package was used to provide the examples shown herein.
- [0087] b) Linear Discriminant Analysis. This involves finding weighted combinations of the features that give the best separation of the classes. These methods work well with correlated data, but not in data with similar means and different variances. Several statistical packages were used (SPSS, SAS and R), depending on the performance estimates and graphical outputs required. Fisher's linear discriminant function was used to obtain the classifier that minimized the error rate. A canonical discriminant function was used to compute receiver operating characteristic (ROC) curves showing the trade-off between sensitivity and selectivity as the decision threshold is changed.
- [0088] c) Logistic Regression. This is a non-linear transformation of the linear regression model: the dependent variable is replaced by a log odds ratio (logit). Linear regression, like discriminant analysis, belongs to a class of statistical methods founded on linear models. Such models are based on linear relationships between the explanatory variables.
- [0089] With a sufficient number of samples it is possible, using the above techniques and software packages, to search for combinations of features giving good discrimination between the classes. Other exemplary statistical analysis and/or pattern recognition methods are the linear

Discriminant Function Method in SPSS and Logistic Regression Method in R and SAS. SPSS is the full product name and is available from SPSS, Inc., located at SPSS, Inc. Headquarters, 233 S. Wacker Drive, 11th floor, Chicago, Illinois 60606 (www.spss.com). SAS is the full product name and is available from SAS Institute, Inc., 100 SAS Campus Drive, Cary, NC 27513-2414, USA (www.sas.com). R is the full product name and is available as Free Software under the terms of the Free Software Foundation's GNU (General Public License). http://www.r-project.org/.

[0090] In some embodiments, a correlation matrix is obtained. Correlation measures the amount of linear association between a pair of variables. A correlation matrix is obtained by correlating the data obtained with one marker to data obtained with another marker. A threshold correlation number may be set, for example, 50% correlation. In this case, all markers with a correlation number of 50% or higher would be considered correlate markers.

[0091] In some embodiments of the present invention, user supplied weighting factors may be used to obtain optimized panels. Weighting may be related to any factor. For example, certain markers may be weighted higher than others due to cost, commercial considerations, misclassifications or error rates, prevalence of a generic disease state in a geographic location, prevalence of a specific disease state in a geographic location, redundancy and availability of probes. Some factors related to cost that may encourage a user to weight certain markers higher than others is the cost of the probe and commercial access issues, such as license terms and conditions. Some factors related to commercial considerations that may encourage a user to weight certain markers higher than others are Research and Development (R&D) time, R&D cost, R&D risk, i.e., the probability that the probe will work, cost of final analytical instrument, final performance and the time to market. In a detection panel, for example, some factors related to misclassifications or error rates that may encourage a user to weight some markers higher than others is that it may be desirable to minimize false negatives. In a discrimination panel, on the other hand, it may be desirable to minimize false positives. Some factors related to prevalence of a generic or specific disease state in a geographic area that may encourage a user to weight some probes higher than others are that in some geographic locations the incidence of certain generic or specific diseases are more or less prevalent. With respect to redundancies, in some instances it is desirable to have redundancies in the panel. For example, if for some reason one probe fails to be detected, due to the biological variability of the markers in the panel, a disease state will still

be detected by the other markers. In some embodiments, markers that are preferred redundant markers may be weighted more heavily.

[0092] The invention is flexible in being adaptable to the availability of features where cost or supply problems may not allow the very best combination. In one embodiment, the invention can simply be applied to the available features to find an alternative combination. In another embodiment, the algorithm is used to select features that allow cost weightings to be included in the selection process to arrive at a minimum cost solution. In the examples, marker performance estimates for combinations selected from all the markers collected or for only a group of commercially preferred probes are shown. The examples also demonstrate how the C4.5 package can be used to down weight certain probes on the basis of their high cost. These probe combinations may not perform as well as the optimum combination, but the performance might be acceptable in circumstances where cost is a significant factor.

[0093] Some of the methods used allow weightings to be applied to the classes. This is available in C4.5 where the tree design can optimize the cost. Also, the Discriminant Function method gives a single parameter output which can be used to give a desired false positive or false negative probability. A plot of these parameters for different threshold settings is known as the receiver operating characteristic (ROC) curve. An ROC curve shows the estimated percentage of false positive against true positive scores for different threshold levels of a classifier.

[0094] Given the heterogeneous nature of many generic disease states, the panels may be constructed with a degree of redundancy to ensure that the tests have sufficient sensitivity, specificity, positive predictive value (Positive Predictive Value = True Positives/(True Positives + False Positives) and negative predictive value (Negative Predictive Value = True negatives/(False Negatives + True Negatives) to justify their use as a population-based screen. However, local and regional differences may dictate specific use of the tests in different segments of the global market, and so may significantly influence the criteria used to construct the final panel test for a given market. While the optimization of clinical utility is of utmost importance, local factors including affordability (cost), technical competence, laboratory and healthcare provider resources, workflow issues, manpower requirements, and availability of the probes and labels will contribute to a final, local selection of the markers used in the panel. Well known linear discriminant function analysis is used to include and assess all potential selection factors, by which each local factor is represented by a term in the equation, and each is weighted

according to its locally determined significance. In this way, a panel test optimized for use in one world region may differ from a panel test optimized for use in a different region.

[0095] Once detection or discrimination panels have been designed using the above described method, the next step is to validate the panel using known cytology samples. Prior to validation, optional optimization steps may be performed. In some embodiments, the method for collecting cytology samples may be improved. This encompasses methods of obtaining the sample from the patient as well as methods for mixing the cytology sample. In other embodiments, the cytology presentation methods may be improved. For example, identifying optimal fixatives (preservation fluids) or transportation fluids.

[0096] The cytology samples used to validate the panels produced using the gold standard histology samples are cytology samples with known diagnoses. These samples may be collected using any method known by those of skill in the art. For example, sputum samples can be collected by spontaneous production, induced production and through the use of agents that enhance sputum production. The sample is contacted with each probe in the panel and the level and pattern of binding of the probes is analyzed to determine the performance of the panel. In some embodiments, it may be necessary to further optimize the panel. For example, it may be necessary to remove a probe from the panel. Or, it may be necessary to add an additional probe to the panel. Additionally, it may be necessary to replace one probe on the panel with another probe. If a new probe is added, this probe may be a correlate marker as determined from a correlation matrix. Alternatively, the probe may be a functionally similar marker. Once the panel is optimized, the panel may proceed for further testing in clinical studies.

[0097] In other embodiments, it is not necessary to optimize the panel. If the results with the cytology samples correlate with the results from the histology samples, there may not be a need to optimize the panel and the panel may proceed for further testing in clinical studies.

4. Methods of Use

[0098] Once a panel is obtained using the above described method, it may be applied to cytologic samples. To illustrate the method, cancer, especially lung cancer, will be exemplified. Similar steps and procedures will be applied for other disease states. It is to be expected that cells shed from the specified lesions will stain in a similar fashion and show in a cytologic sample, such as a fine need aspiration, sputum, urine, in a similar fashion as in the histologic pathology samples used to obtain the panel.

[0099] The basic method of the present invention typically involves two steps. First, a cytological sample suspected of containing diseased cells is contacted with a panel containing a plurality of agents, each of which quantitatively binds to a disease marker. Then, the level or pattern of binding of each agent to a disease marker is detected. The results of the detection may be used to diagnose the presence of a generic disease or to discriminate among specific disease states. An optional preliminary step is identifying an optimized panel of agents that will aid in the detection of a disease or the discrimination between disease states in a cytologic sample. [0100] Cytology specimens may include, but are not limited to, cellular samples collected from body fluids, such as blood, urine, spinal fluids, and lymphatic systems; epithelial cell-based organ systems, such as the pulmonary tract, e.g., lung sputum, urinary tract, e.g., bladder washings, genital tract, e.g., cervical Pap smears, and gastrointestinal tract, e.g., colonic washings; and fine needle aspirations from solid tissue sites in organs and systems such as the breast, pancreas, liver, kidneys, thyroid, bone marrow, muscles, prostate, and lungs; biopsies from solid tissue sites in organs and systems such as the breast, pancreas, liver, kidneys, thyroid, bone marrow, muscles, prostate, and lungs; and histology specimens, such as tissue from surgical biopsies.

[0101] An illustrative panel of agents according to the present invention includes any number of agents that allows for accurate detection of malignant cells in a cytological sample. Molecular markers envisioned by the present invention may be any molecule that aids in the detection of malignant cells. Markers may be selected for inclusion in a panel based on several different criteria relating to changes in level or pattern of expression of the marker. Preferred are molecular markers where changes in expression: occur early in tumor progression, are exhibited by a majority of tumor cells, allow for detection of in excess of 75% of a given tumor type, most preferably in excess of 90% of a given tumor type and/or allow for the discrimination between histologic types of cancer.

[0102] The first step of the basic method is the detection of changes in the level or pattern of expression of the panel of agents in a cytological sample. This step typically involves contacting the cytologic sample with an agent, such as a labeled polyclonal or monoclonal antibody or fragment thereof or a nucleic acid probe, and observing the signal in individual cells. Detection of cells where there is a change in signal is indicative of a change in the level of expression of the molecular marker to which the label probe is directed. The changes are based on an increase or

decrease in the level of expression relative to nonmalignant cells obtained from the tissue or site being examined.

[0103] An analysis of the changes in the level or pattern of expression of a panel of agents enables a skilled artisan to determine, with high sensitivity and high specificity, whether malignant cells are present in the cytologic sample. The term "sensitivity" refers to the conditional probability that a person having a disease will be correctly identified by a clinical test, (the number of true positive results divided by the number of true positive and false negative results). Therefore, if a cancer detection method has high sensitivity, the percentage of cancers detected is high e.g., 80%, preferably greater than 90%. The term "specificity" refers to the conditional probability that a person not having a disease will be correctly identified by a clinical test, (i.e., the number of true negative results divided by the number of true negative and false positive results). Therefore, if a cancer detection method has high specificity, 80%, preferably 90%, more preferably 95%, the percentage of false positives the method produces is low. A "cytologic sample" encompasses any sample collected from a patient that contains that patient's cells. Examples of cytological samples envisioned by the present invention include body fluids, epithelial cell-based organ system washings, scrapings, brushings, smears or effusions, and fine-needle aspirates and biopsies.

[0104] Use of the markers described in this invention assumes that it is possible to obtain an adequate cytologic sample routinely and that the samples can be adequately preserved for subsequent evaluation. The cytologic sample may be processed and stored in a suitable preservative. Preferably, the cytologic sample is collected in a vial containing the preservative. The preservative is any molecule or combination of molecules known to maintain cellular morphology and inhibit or block degradation of cellular proteins and nucleic acids. To ensure proper fixation, the sample may be mixed at the collection site at high speeds to disaggregate the sample and/or break up obscuring material such as mucus, thereby exposing the cells to the preservative.

[0105] Once a specimen is obtained, it is desirable to homogenize it, using an appropriate mixing device. This permits using aliquots for multiple purposes, including the possibility of sending aliquots to more than one testing site, as well as preparing multiple slides and/or multiple depositions on a slide. The initial homogenization of the specimen and of each aliquot before use

will ensure that each individual slide will have substantially the same distribution of cells, so that comparisons of results from one slide to another will be meaningful.

[0106] Preparation of a specimen for analysis involves applying a sample to a microscope slide using methods including, but not limited to, smears, centrifugation, or deposition of a monolayer of cells. Such methods may be manual, semi-automated, or fully automated. The cell suspension may be aspirated depositing the cells on a filter and a monolayer of cells transferred to a prepared slide that may be processed for further evaluation. By repeating this process additional slides may be prepared as necessary. The present invention encompasses detection of one molecular marker per slide. Detection of several molecular markers per slide is also envisioned.

Preferably, 1-6 markers are detected per slide. In some embodiments 2 markers are detected per

Preferably, 1-6 markers are detected per slide. In some embodiments 2 markers are detected per slide. In other embodiments, 3 markers are detected per slide.

[0107] The present invention contemplates detecting changes in molecular marker expression at the DNA, RNA or protein level using any of a number of methods available to an ordinary skilled artisan. Detection of the changes in the level or pattern of expression of the molecular markers in a cytologic sample generally involves contacting a cytologic sample with a polyclonal or monoclonal antibody or fragment thereof or a nucleic acid sequence that is complementary to the nucleic acid sequence encoding a molecular marker in the panel, collectively "probes", and a label. Typically, the probe and label components are operatively linked so that when the probe reacts with the molecular marker a signal is emitted (a "labeled probe"). Labels envisioned by the present invention are any labels that emit or enable a signal and allow for identification of a component in a sample. Preferred labels include radioactive, fluorogenic, chromogenic or enzymatic moieties. Therefore, possible methods of detection include, but are not limited to, immunocytochemistry; proteomics, such as immunochemistry; cytogenetics, such as in situ hybridization, and fluorescence in situ hybridization; radiodetection, cytometry and field effects, such as MACs and DNA ploidy (the quantitation of stoichiometrically-stained nuclear DNA using automated computerized cytometry) and; cytopathology, such as quantitative cytopathology based on morphology. The signal generated by the labeled probe is preferably of sufficient intensity to permit detection by a medical practitioner or technician.

[0108] Once the slide is prepared, a medical practitioner conducts a microscopic review of the slides in order to identify cells that exhibit a change in marker expression characteristic of a diagnosis of cancer. The medical practitioner may use an image analysis system and automated

microscope to identify cells of interest. Analysis of the data may make use of an information management system and algorithms that will assist the physician in making a definitive diagnosis and select the optimal therapeutic approach. A medical practitioner may also examine the sample using an instrument platform that is capable of detecting the presence of the labeled agent.

- [0109] A molecular diagnostic panel assay will result in one or more glass microscope slides with labeled cells and/or tissue sections. The challenge for human experts to assess these (cyto)pathology multilabeled-cell preparations objectively and with clinically meaningful results is a virtually insurmountable detection and perception problem for any human being.
- [0110] Computer-aided imaging systems (i.e., Photonic MicroscopesTM) can be developed and used to assess quantitatively and reproducibly the amount and location of probe-labeled cells and tissues. Such Photonic MicroscopesTM combine robotic slide-handling capabilities, data management systems (e.g., medical informatics), and quantitative digital (optical and electronic) image analysis hardware and software modules to detect and report cell-based probe content and localization data that cannot be obtained by human visualization with comparable sensitivity and accuracy. These probe data can be used to characterize and differentiate cellular samples based upon their related characteristics and differences in their respective cell-based markers for a variety of disease states.
- [0111] The present methodology is a methodology whereby the molecular diagnostic panels are applied to cell-based specimens and samples, and whereby computer-aided imaging systems are subsequently used to quantify and report the results of the molecular diagnostic panel tests. Such imaging systems can be used to evaluate cell-based samples in which multiple probes are used simultaneously on a given slide-based sample, and in which the probes can be separately analyzed, quantified, and reported because the probes are differentiated by color on the microscope cytology or histology slide.
- [0112] The signals generated by a labeled agent in the sample may, if they are of appropriate type and of sufficient intensity, be detected by a human reviewer (e.g., pathologist) using a standard microscope or a Computer-Aided Microscope [167]. The Computer-Aided Microscope is an ergonomic, computer-interfaced microscope workstation that integrates mouse-driven control of microscope operation (e.g., stage movement, focusing) with computerized automation of key functions (e.g., slide scanning patterns). A centralized Data Management System stores,

organizes and displays relevant patient information as well as results from all specimen screenings and pathologist reviews. An identification number that is imprinted onto barcodes and affixed to each sample slide uniquely identifies each sample in the database, and relates it to the original specimen and the patient.

[0113] In a preferred embodiment the signals generated by a labeled agent in the sample will be detected and quantitated using an automated image analysis system, or Photonic Microscope, interfaced to the centralized Data Management System. The Photonic Microscope provides fully automated software control of the microscope operations and incorporates detectors and other components appropriate for quantitation even of signals not detectable by human reviewers, such as very faint signals or signals from radiolabeled moieties. The location of detected signals is stored electronically for rapid relocation by automated instruments, and for human review using a Computer-Aided Microscope [168].

[0114] The centralized Data Management System archives all patient and sample data using the bar-coded identification number. The data may be acquired asynchronously, from a multiplicity of sites, and may be derived from multiple reviews and analyses by human cytologists and/or automated analyzers. These data may include results from multiple sample slides representing aliquots from a single previously homogenized patient specimen. Part or all of the data may be transferred to or from a hospital's Laboratory Information System to meet reporting, archiving, billing or regulatory requirements. A single, comprehensive report with integrated results from panel tests and human reviews may be generated and delivered to the physician in hardcopy, or electronically through networked computers or the Internet.

[0115] In some embodiments, the instant method allows for differential discrimination of different diseases, such as different histologic types of cancers. The term "histologic type" refers to specific disease states. Depending on the general disease state there can be one or several histologic types. For example, lung cancer includes, but is not limited to, squamous cell carcinoma, adenocarcinoma, large cell carcinoma, small cell carcinoma and mesothelioma. Knowledge of the histologic type of cancer affecting a patient is very useful because it helps the medical practitioner to localize and characterize the disease and to determine the optical treatment strategy.

[0116] In order to determine the specific disease state, a panel of markers is selected that allows for discrimination between specific disease states. For example, within a panel of molecular

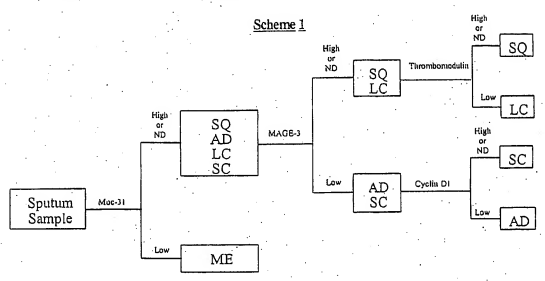
markers, a pattern of expression may be identified that is indicative of a particular histologic type of cancer. The detection of the level of expression of the panel of molecular markers is achieved by the above-described methods. Preferably, a panel of 1-20 molecular markers is employed to discriminate among the various histologic types of lung cancer. However, most preferably, 4-7 markers are used. Decision trees may be developed to aid in discriminating between different histologic types based on patterns of marker expression.

[0117] In addition to allowing for the detection of malignant cells in a cytologic sample, the instant invention has utility in the molecular characterization of the disease state. Such information is often of prognostic significance and can assist the physician in the selection of the optimal therapeutic approach for a particular patient. In addition, the panel of markers described in this invention may have utility in monitoring the patient for either recurrence or to measure the efficacy of the therapy being used to treat the disease.

[0118] By way of non-limiting example, the presence of lung cancer may be detected by a lung cancer detection panel and the specific type of lung cancer may be detected by a discrimination panel. If the medical practitioner determines that malignant cells are present in the cytologic sample, a further analysis of the histologic type of lung cancer may be performed. The histologic type of lung cancer encompassed by the present invention includes but is not limited to squamous cell carcinoma, adenocarcinoma, large cell carcinoma, small cell carcinoma and mesothelioma. Figure 1 illustrates molecular markers that are preferable markers to be included in a panel for identifying different histologic types of lung cancer. The column labeled "%" indicates the percentage of tumor specimens that express a particular marker.

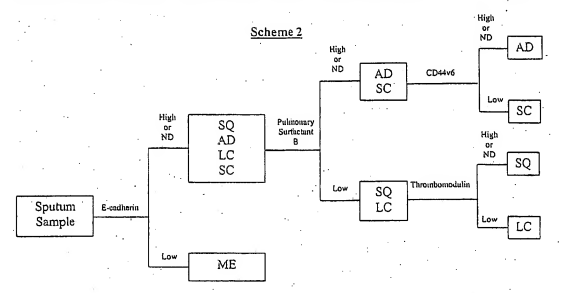
[0119] In determining the various histologic types of lung cancer, the relative level of expression of a marker is analyzed. Figure 2 illustrates how different markers may be used to discriminate among different histologic types of cancer. In this table, SQ indicates squamous cell carcinoma, AD indicates adenocarcinoma, LC indicates large cell carcinoma, SC indicates small cell carcinoma and ME indicates mesothelioma. The numbers appearing in each cell represent frequency of marker change in one cell type versus another. To be included in the table, the ratio must be greater than 2.0 or less than 0.5. A number larger than 100 generally indicates that the second marker is not expressed. In such cases the denominator was set at 0.1 for the purpose of the analysis. Finally, empty cells represent either no difference in expression or the absence of expression data.

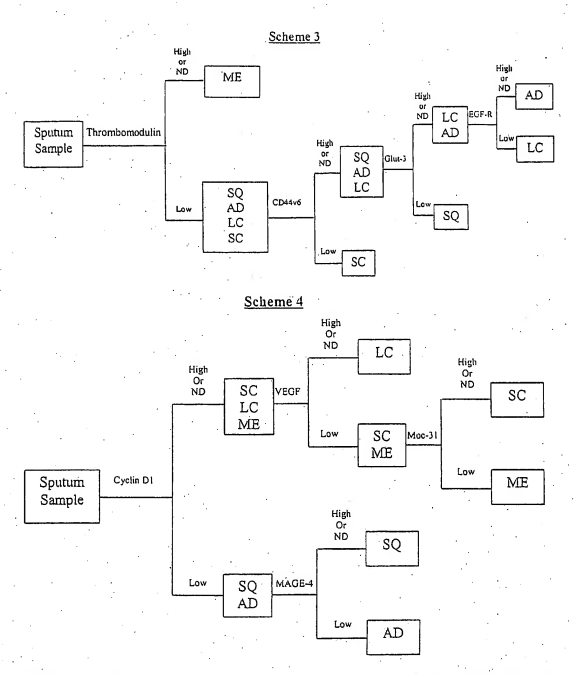
One method for analyzing the data collected is to construct decision trees. Schemes 1-4 are examples of decision trees that may be constructed to enable a differential determination of a histologic type of lung cancer using the patterns of expression. The present invention is in no way limited to the decision trees presented in Schemes 1-4. The relative level of expression of a marker can be higher, lower, or the same (ND) as the level of expression of the molecular marker in a malignant cell of a different histologic type. Each scheme enables a distinction between five histologic types of lung cancer through the use of the indicated panel of molecular markers. [0121] For example, in Scheme 1 the panel consists of HERA, MAGE-3, Thrombomodulin and Cyclin D1. First the sample is contacted with a labeled probe directed toward HERA. If the expression of HERA is lower than the control, the test indicates that the histologic type of lung cancer is mesothelioma (ME). If, however, the expression is higher or the same as the control, the sample is contacted with a probe directed toward MAGE-3. If the expression of MAGE-3 is lower than the control, the sample is contacted with a labeled probe directed toward Cyclin D1 and a determination of small cell carcinoma (SC) or adenocarcinoma (AD) is possible. If the expression of MAGE-3 is higher than or the same as the control, the sample is contacted with a labeled probe directed toward Thrombomodulin and a determination of squamous cell carcinoma (SC) or large cell carcinoma (LC) is possible.



[0122] In Scheme 2 the panel consists of E-Cadherin, Pulmonary Surfactant B and Thrombomodulin. First the sample is contacted with a labeled probe directed toward E-Cadherin. If the expression of E-Cadherin is lower than the control, the test indicates that the

histologic type of lung cancer is mesothelioma (ME). If, however, the expression is higher or the same as the control, the sample is contacted with a probe directed toward Pulmonary Surfactant B. If the expression of Pulmonary Surfactant B is lower than the control, the sample is contacted with a labeled probe directed toward Thrombomodulin and a determination of squamous cell carcinoma (SQ) or large cell carcinoma (LC) is possible. If the expression of Pulmonary Surfactant B is higher than or the same as the control, the sample is contacted with a labeled probe directed toward CD44v6 and a determination of adenocarcinoma (AD) and small cell carcinoma (SC) is possible. (See Schemes 3 and 4 for more examples of decision trees).





- [0123] A preferred method involves using panels of molecular markers where differences in the pattern of expression permits the discrimination between the various histologic type of lung cancer.
- [0124] Many different decision trees may be constructed to analyze the patterns of marker expression. This information may be used by physicians or other healthcare providers to make patient management decisions and select an optimal treatment strategy.

5. Reporting of Results of Panel Analysis

[0125] The results from the panel analysis may be reported in several ways. For example, the results may be reported as a simple "yes or no" result. Alternatively, the result may be reported as a probability that the test results are correct. For example, the results from a detection panel study may indicate whether a patient has a generic disease state or not. As the panel also reports the specificity and sensitivity, the results may also be reported as the probability that the patient has a generic disease state. The results from a discrimination panel analysis will discriminate among specific disease states. The results may be reported as a "yes or no" with respect to whether the specific disease state is present. Alternatively, the results may be reported as a probability that a specific disease state is present. It is also possible to perform several discrimination panel analyses on a specimen from one patient and report a profile of the probabilities that the disease state present is a specific disease state with respect to the other possibilities. The other possibilities may also include false positives.

[0126] In embodiments in which a profile of the probabilities of each specific disease state being present is produced, there are several possible outcomes. For example, it is possible that all of the probabilities will be a very small probability. In this instance, it is possible that the doctor will conclude that the patient's specimen diagnosis is a false positive. It is also possible that all of the probabilities will be low except for one that is above 80-90%. In this instance, it is possible that the doctor will conclude that the test verifies that the patient has the specific disease state that indicated the high probability. It is also possible that most of the probabilities will be low, but similarly high probabilities are reported for two specific disease states. In this case, a doctor may recommend more extensive panel testing to ensure that the correct disease state is identified. Another possibility is that all of the probabilities reported will be low, with one being slightly higher than the rest but not high enough to be in the 80-90% range. In this case, a doctor may recommend more extensive panel testing to ensure that the correct disease state is identified and/or to rule out metastatic cancer from a remote primary tumor of a different cancer type.

[0127] The following Example is illustrative of the method of the invention for selecting a disease detection panel, disease discrimination panels, validation of the panels and use of the panels in the clinic to screen for a disease and to discriminate among different subtypes of the disease. Lung cancer was selected for this illustrative example, in part because of its importance to world health, but it will be appreciated that similar procedures will apply to other types of

cancer, as well as to infectious, degenerative and autoimmune diseases, according to the foregoing general disclosure.

ILLUSTRATIVE EXAMPLES.

I. Lung Cancer

[0128] The present method was used to develop lung cancer detection panels as well as single lung cancer type specific discrimination panels. Lung cancer is an extremely complex collection of diseases that can be segregated into two main classes. Non-small cell lung carcinoma (NSCLC) that accounts for approximately 70 to 80% of all lung cancers can be further subdivided into three main histologic types including squamous cell carcinoma, adenocarcinoma, and large cell carcinoma. The remaining 20 to 30% of lung cancer patients present with small cell lung carcinoma (SCLC). In addition, malignant mesothelioma of the pleural space, can develop in individuals exposed to asbestos and will often spread widely invading other thoracic structures. Different forms of lung cancer tend to localize in different regions of the lung, have different prognoses, and respond differently to various forms of therapy.

[0129] According to the latest statistics from the World Health Organization (Globocan 2000), lung cancer has become the most common fatal malignancy in both men and women with an estimated 1.24 million new cases and 1.1 million deaths each year. In the U.S. alone, the National Cancer Institute reports that there are approximately 186,000 new cases of lung cancer and each year 162,000 people die of the disease, accounting for 25% of all cancer-related deaths. In the U.S., overall 1-year survival for patients with lung cancer is 40%, however, only 14% live 5 years. In other parts of the world, 5-year survival is significantly lower (5% in the UK). The high mortality of lung cancer can be attributed to the fact that most patients (85%) are diagnosed with advanced disease when treatment options are limited and the disease is likely to have metastasized. In these patients, 5-year survival is between 2-30% depending of the stage at the time of diagnosis. This is in sharp contrast to cases where patients are diagnosed early and 5year survival is greater than 75%. While it is true that a number of new chemotherapeutic agents have been introduced into clinical practice for the treatment of advanced lung cancer, to date, none have yielded a significant improvement in long-term survival. Even though patients with early stage disease can presumably be cured by surgery, they remain at significant risk, as there is a high probability that they will develop a second malignancy. Thus, for the lung cancer patient, early detection and treatment followed by aggressive monitoring provides the best chance of

achieving significant improvements in long-term survival along with a reduction in morbidity and cost.

[0130] At the present time, a patient is suspected of having lung cancer either because of a suspicious lesion on X-ray or because the patient becomes symptomatic. As a result, most patients are diagnosed with relatively late stage disease. In addition, because most methods lack sufficient sensitivity with respect to the detection of early stage disease, the current policy of the U.S. National Cancer Institute (NCI), National Institutes of Health, recommends against screening for lung cancer even in populations of patients who are at significant risk. In this embodiment of the present invention, however, sputum cytology is employed to provide a relatively noninvasive, more effective and cost-effective means for the early detection of lung cancer.

[0131] The specificity of sputum cytology is relatively high. Recent studies have indicated that experienced cytotechnologists are able to recognize malignant or severely dysplastic cells with a high degree of accuracy and reliability [10]. While the detection rate can be as high as 80 to 90% when samples are collected from patients with a relatively advanced disease [11,12], overall, sputum cytology has a sensitivity of only 30-40% [13,14]. The low sensitivity of sputum cytology is particularly important given that obtaining and preparing the specimen can be relatively expensive. Furthermore, failing to detect a malignancy can significantly delay treatment thereby reducing the chance of achieving a cure.

[0132] The selection of an "at-risk" population can also influence the value of sputum cytology as a screening tool. Individuals who are at significant risk include those with a prior diagnosis of lung cancer, long-term smokers or former smokers (>30 pack years) and individuals with long-term exposure to asbestos or pulmonary carcinogens. People with a genetic predisposition or familial history are also included in an "at-risk" population. Such individuals are likely to benefit from testing. While the inclusion of individuals with lower risk may result in an increase in the absolute number of cases detected, it would be hard to justify the substantial increase in healthcare costs.

[0133] Other factors that contribute to the relatively poor performance of conventional sputum cytology include the location of the lesion, tumor size, histologic type, and the quality of the sample. Squamous-cell carcinoma accounts for 31% of all primary pulmonary neoplasms. Most of these tumors arise from segmental bronchi and extend to the proximal lobar and distal

subsegmental branches [15]. For this reason, sputum cytology is reasonably effective (79%) in detecting these lesions. Currently, squamous cell carcinoma is viewed as the only type of lung cancer that is amenable to cytologic detection in an *in situ* and radiologically occult stage [15], as sloughed cells are more likely to be available for evaluation. In one large study where patients were followed with both chest X-ray and sputum cytology, 23% of all lung cancers were detected by cytology alone, suggesting that the tumors were early stage and radiologically occult [16]. In another study [17], sputum cytology detected 76% of patients with radiologically occult tumors. [0134] In the case of adenocarcinoma, 70% of tumors occur in the periphery of the lung making it less likely that malignant cells will be found in a conventional sputum specimen. For this reason, adenocarcinomas are rarely detected by sputum cytology (45%) [12;18,19], an important consideration, since the incidence of adenocarcinoma appears to be increasing, particularly in women [20-22].

- [0135] Tumor size can also affect the likelihood of achieving a correct diagnosis, a factor that is particularly important when considering a screening test for the detection of disease in asymptomatic individuals. While there is only a 50% chance that tumors <24mm will be read as a true positive, the probability of detecting a larger lesion is in excess of 84% [12].
- [0136] Recent reports also indicate that the cellularity of the specimen will affect the sensitivity of sputum cytology [14,23]. In general, patients with squamous cell carcinoma produce specimens with significant numbers of tumor cells, thereby increasing the likelihood of a correct diagnosis [14,23]. For patients with adenocarcinoma, the presence of tumor cells in a sputum specimen is reported to be less than 10% in 95% of the specimens and less than 2% in 75% of specimens, making the diagnosis significantly more difficult.
- [0137] The degree of differentiation can also influence the ability of a pathologist to detect malignant cells, particularly in cases of adenocarcinoma. Well-differentiated tumor cells frequently resemble nonneoplastic respiratory epithelial cells. In the case of small-cell lung carcinoma, sputum samples often contain nests of loosely aggregated cells that have a distinct appearance. However, techniques currently used to process sputum samples tend to disaggregate the cells, making a diagnosis more difficult.
- [0138] Sample quality is another factor that can contribute to the low sensitivity of sputum cytology. Recent reports suggest that it is possible to obtain adequate samples from 70-85% of subjects. However, achieving this measure of success often requires that patients provide

multiple specimens [13]. This procedure is inconvenient, time-consuming and costly. Patient compliance is also generally low, as patients are frequently asked to collect over several days [13]. Of equal importance is the observation that former smokers, while at significant risk for developing lung cancer, often fail to produce an adequate specimen. Sample preservation and processing is another critical factor that can affect the value of sputum cytology as a diagnostic test.

[0139] Lastly, even if adequate samples could be obtained and optimally prepared, cytotechnologists generally still have to review 2-4 slides per specimen, each typically taking up to four minutes [24]. Given the low sensitivity, high technical complexity and labor intensity of conventional sputum cytology, it is not surprising that this test has been almost universally rejected as a population-based screen for the early detection of lung cancer [25].

[0140] Even if these technical issues were resolved, the low sensitivity of sputum cytology remains a significant problem. The high incidence of false negative results can significantly delay the patient receiving potentially curative therapy. While it may be possible to develop tests with greater sensitivity, such improvements must not come at the cost of specificity. An increase in the number of false positive results would subject patients to unnecessary, often invasive and costly, follow-up and would have a negative impact on the patient's quality of life. The present invention overcomes many of the limitations associated with previous methods of early cancer detection, including those related to the use of sputum cytology for the early detection of lung

[0141] Lung cancer is a heterogeneous collection of diseases. To ensure that a test has the necessary level of sensitivity and specificity to justify its use as a population based screen, the present invention envisions using, for example, a library of 10 to 30 cellular markers to develop panels. Selection of the library of this invention was based on a review and reanalysis of the relevant scientific literature where, in most cases, marker expression was measured in biopsy specimens taken from patients with lung cancer in an attempt to link expression with prognosis. [0142] For example, a preferred panel for early detection, characterization, and/or monitoring of lung cancer in a patient's sputum may include molecular markers for which a change in expression occurred in at least 75% of tumor specimens. An exemplary panel includes markers selected from VEGF, Thrombomodulin, CD44v6, SP-A, Rb, E-Cadherin, cyclin A, nm23, telomerase, Ki-67, cyclin D1, PCNA, MAGE-1, Mucin, SP-B, HERA, FGF-2, C-MET, thyroid

cancer.

transcription factor, Bcl-2, N-Cadherin, EGFR, Glut-1, ER-related (p29), MAGE-3 and Glut-3. A most preferred panel includes molecular markers for which a change in expression occurs in more than 85% of tumor specimens. An exemplary panel includes molecular markers selected from Glut1, HERA, Muc-1, Telomerase, VEGF, HGF, FGF, E-cadherin, Cyclin A, EGF Receptor, Bcl-2, Cyclin D1 and N-cadherin. With the exception of Rb and E-cadherin, a diagnosis of lung cancer is associated with an increase in marker expression. A brief description of the library of probes/markers utilized in the present example is provided below in Table 4. It is noted that the numbering of the antibodies in the table below is consistent with the number of the antibodies/probes/markers throughout this example.

Table 4:

		Probes and Markers fo	r Lung Panel
No.		Full Name of Antibody Probe	Target Marker Name/Description
1	VEGF	anti-VEGP	Vascular Endotnelial Growth Factor protein
2	Thrombomodulin	anti-Thrombomodulin	trams-membrane glycoprotein
3	CD44v6	anti-CO44v6	cell surface glycoprotein (CD44 vanant 6 gene); cell achesion molecule
4	SP-A	anti-Surfactant Apoprotein A	pulmonary surfaciant apoprotein
5	Retinoblastoma	anti-Retinoblastoma gene product	phosphoprotein
- 6	E-Cadherin	anti-E-Cadherin	transmembrane Ca dependent cell achesion molecule
7	Cyclin A	anti-Cyclin A	protein subunit of cyclin-dependent kinase enzymes; for cell cycle regulation
8	nm23	lanti-nm23	2 closely related proteins produced by nm23-H1 and -H2 genes
9	Telomerase	anti-Telomerase	Inbonucieoprotein enzyme for chromosome repair
10	Mib-1 (KG-67)	lanti-KI-87	nuclear protein; expressed in proliferating cells
11	Cyclin D1	anti-Cyclin D1	protein subunit of cyclin-dependent kinase enzymes; for cell cycle regulat
12	PCNA	anti-Proliferating Cell Nuclear Antigen	protein cofactor for DNA polymerase delta
13	MAGE-1	anti-Melanoma-Associated Antigen 1	cell recognition protein coded by MAGE family of genes
14	Mucin 1 (MUC-1)	anti-Mucin 1	cell surface and secreted mucin (highly glycosylated protein)
15	SP-B	anti-mature Surfactant Apoprotein 8	pulmonary surfactant apoprotein
16	HERA .	anti-Human Epithelial Related Antigen (MOC-31)	cell surface antigen (transmemorane protein)
17	FGF-2 (basic FGF)	anti-Fibroplast Growth Factor	protein that binds to cell surface
18	ic-MET	anti-c-MET .	trans-memorane receptor protein for Hepatocyte Growth Factor (HGF)
19	Thyroid Transcription Factor 1	anti-TTF-1	regulator of thyroid-specific genes; also expressed in lung
20	BCL-2	anti-BCL2	intracellular membrane-bound protein encoded by BCL2 gene
21	P120	lanti-p120	Proliferation-Associated Nucleolar Antigen protein
22	N-Cacherin	anti-N-Cadherin	transmembrane Ca dependent cell adhesion molecule :
23	EGFR	anti-EGFR	Epidermal Growth Factor Receptor; transmembrane glycoprotein
24	Glut 1	anti-Glut 1	Glucose-transporting, transmembrane Glut family of proteins
25	ER-related (p29)	anti-ER-related P29; anti-HSP 27	Estrogen Receptor-related p29 protein; Heat Shock protein 27
26	Mage 3	anti-Melanoma-Associated Antigen 3	cell recognition protein coded by MAGE family of genes
27	Glut 3	anti-Glut 3	Glucose-transporting, transmemorane Glut family of proteins
28	PCNA (higher dilution)	anti-Proliferating Cell Nuclear Antigen	protein collector for DNA polymerase delta

[0143] Each molecular marker in the preferred panel is described below. Table 5, reciting the percentage of expression of the markers in tissue for each type of lung cancer is provided at the end of this section.

Glucose Transporter Proteins (Glut 1 and Glut 3) [26-28]

[0144] Glucose Transporter-1 (Glut 1) and Glucose Transporter-3 (Glut-3) are a ubiquitously expressed high affinity glucose transporter. Tumor cells often display higher rates of respiration, glucose uptake, and glucose metabolism than do normal cells, and the elevated uptake of glucose in tumor cells is thought to be mediated by glucose transporters. Overexpression of certain types

of GLUT isoforms has been reported in lung cancer. The cellular localization of Glut 1 is in the cell membrane. GLUT-1 and GLUT-3 are disease markers useful for detection of a disease state. [0145] Malignant cells exhibit an increase in glucose uptake that appears to be mediated by a family of glucose transporter proteins (Gluts). Oncogenes and growth factors appear to regulate the expression of these proteins as well as their activities. Members of the Glut family of proteins exhibit different patterns of distribution in various human tissues and rapid proliferation is often associated with their overexpression. Recent evidence suggests that Glut1 is expressed by a large percentage of NSCLC and by a majority of SCLC.

[0146] While the expression of Glut 3 is relatively low in both NSCLC and SCLC a significant percentage (39.5%) of large cell carcinomas express the protein. In stage I tumors, 83% express Glut1 at some level with 75-100% of cells staining in 25% of cases. These data would suggest that Glut1 overexpression is a relatively early event in tumor progression. Glut1 immunoreactivity has also been detected in > 90% of stage II and IIIA cancers. There also appears to be an inverse correlation between Glut1 and Glut3 immunoreactivity and tumor differentiation. Tumors expressing high levels of Glut1 appear to be particularly aggressive that are associate with a poor prognosis. In cases were tumors were negative for the proteins better survival was observed.

Human Epithelial Related Antigen (HERA) [29,30]

[0147] HERA is a transmembrane glycoprotein with an, as yet, unknown function. HERA is present on most normal and malignant epithelia. Recent reports suggest that the while HERA expression is high in all histologic types of NSCLC making it useful as a detection marker. In contrast HERA expression is absent in mesothelioma and thus suggesting would have utility as a discrimination marker. The cellular localization of HERA is the cell surface.

Basic Fibroblast Growth Factor (FGF) [31-34]

[0148] Basic Fibroblast Growth Factor (FGF) is a polypeptide growth factor with a high affinity for heparin and other glycosaminoglycans. In cancer, FGF functions as a potent mitogen, plays a role in angiogenesis, differentiation, and proliferation, and is involved in tumor progression and metastasis. FGF overexpression frequently occurs in both SCLC and squamous cell carcinoma. In many cases (62%), the cells also express the FGF receptor suggesting the presence of an autocrine loop. Forty-eight percent of Stage 1 tumors overexpress FGF. The frequency of FGF in Stage II lung cancer is 84%. Expression of either the growth factor or its

receptor was associated with the poor prognosis. Five-year survival rates for those patients with stage I disease were 73% for those expressing FGF versus 80% for those who were FGF negative. The cellular localization is the cell membrane.

Telomerase [35-42]

[0149] Telomerase is a ribonucleoprotein enzyme that extends and maintains telomeres of eukaryotic chromosomes. It consists of a catalytic protein subunit with reverse transcriptase activity and an RNA subunit with reverse transcriptase activity and an RNA subunit that serves as the template for telomere extension. Cells that do not express telomerase have successively-shortened telomeres with each cell division, which ultimately leads to chromosomal instability, aging and cell death. The cellular localization of telomerase is nuclear.

[0150] Expression of telomerase appears to occur in immortalized cells and enzyme activity is a common feature of the malignant phenotype. Approximately 80-94% of lung tumors exhibit high levels of telomerase activity. In addition, 71% of hyperplasia, 80% of metaplasia, and 82% of dysplasia express enzyme activity. All the carcinoma in situ (CIS) specimens exhibit enzyme activity. The low levels of expression in premalignant tissues is probably related to the fact that only a small percentage of cells (5 and 20%) in the sample express enzyme activity. This is in contrast to tumors where 20-60% of cells may express enzyme activity. Based on a limited number of samples it would appear expression of telomerase activity is also common in SCLC.

Proliferating Cell Nuclear Antigen (PCNA) [43-51]

[0151] PCNA functions as a cofactor for DNA polymerase delta. PCNA is expressed in both S phase of the cell cycle and during periods of DNA synthesis associated with DNA repair. PCNA is expressed in proliferating cells in a wide range of normal and malignant tissues. The cellular localization of PCNA is nuclear.

[0152] Expression of PCNA is a common feature of rapidly dividing cells and is detected in 98% of tumors. Immunohistochemical staining is nuclear with moderate to intense staining detected in 83% of NSCLC. Intense PCNA staining was observed in 51% of p53-negative tumors. However, when both PCNA (> 50% of cells staining) and p53 are overexpressed (>10% of cells stained) the prognosis tends to be poorer with a shorter time to progression. Although frequently detected in all stages of lung cancer, intense staining for PCNA is more common in metastatic disease. Thirty-one percent of CIS also overexpress PCNA.

CD44 [51-58]

[0153] CD44v6 is a cell surface glycoprotein that acts as a cellular adhesion molecule. It is expressed on a wide range of normal and malignant cells in epithelial, mesothelial and hematopoietic tissues. The expression of specific CD44 splice variants has been shown to be associated with metastasis and poor prognosis in certain human malignancies. It is expected to be used for detection and discrimination between squamous cell carcinoma and adenocarcinoma. CD44 is a cell adhesion molecule that appears to play a role in tumor invasion and metastasis. Alternative splicing results in the expression of several variant isoforms. CD44 expression is generally lacking in SCLC and is variably expressed in NSCLC. Highest levels of expression occur in squamous cell carcinoma, thus making it valuable in discriminating between tumor types. In non-neoplastic tissue, CD44 staining is observed in bronchial epithelial cells, macrophages, lymphocytes, and alveolar pneumocytes. There was no significant correlation between CD44 expression and tumor stage, recurrence, or survival particularly when overexpression occurs in early stage disease. In metastatic lesions 100% of squamous cell carcinoma and 75% of adenocarcinoma showed strong CD44v6 positivity. These data would tend to indicate that changes in CD44 expression occur relatively late in tumor progression that could limit its value as an early detection marker. Recent findings suggest that the CD44v8-10 variant is expressed by a majority of NSCLC making it a possible candidate marker.

Cyclin A [59-62]

[0154] Cyclin A is a regulatory subunit of the cyclin-dependent kinases (CDK's) which control the transition points at specific phases of the cell cycle. It is detectable in S phase and during progression into G2 phase. The cellular localization of Cyclin A is nuclear.

[0155] Protein complexes consisting of cyclins and cyclin-dependent kinases function to regulate cell cycle progression. Changes in cyclin expression are associated with genetic alterations affecting the CCDN1 gene. While the cyclins act as regulatory molecules, the cyclin-dependent kinases function as catalytic subunits activating and inactivating Rb.

[0156] Immunohistochemical analysis has revealed that the overexpression of the cyclins is associated with an increase in cellular proliferation as indicated by a high Ki-67 labeling index. Cyclin overexpression occurs in 75% of NSCLC and appears to occur relatively early in tumor progression. Recent reports indicate that 66.7% of stage I/II and 70.9% of stage III tumors overexpress Cyclin A. Nuclear staining is common in poorly differentiated tumors. Expression

of cyclin A is often associated with a decrease in mean survival time and a tendency towards the development of drug resistance. However, increased expression has also been associated with a greater response to doxorubicin.

Cyclin D1 [63-73]

[0157] Cyclin D1, as with Cylcin A, is a regulatory subunit of the cyclin-dependent kinases (CDK's) which control the transition points at specific phases of the cell cycle. Cyclin D1 regulates the entry of cells into S phase of the cell cycle. This gene is frequently amplified and/or its expression deregulated in a wide range of human malignancies. The cellular localization of Cyclin D1 is nuclear.

[0158] Like Cyclin A, cyclin D1 functions to regulate cell cycle progression. Staining of cyclin D1 is predominately cytoplasmic and independent of histologic type. Reports suggest that cyclin D1 overexpression occurs in 40-70% of NSCLC and 80% of SCLC. Cyclin D1, staining was observed in 37.9% of stage I, 60% stage II, and 57.9% of stage III tumors. Cyclin D1 expression has also been seen in dysplastic and hyperplastic tissue providing evidence that these changes occur relatively early in tumor progression. Patients who overexpress cyclin D1 exhibit shorter mean survival time and lower five-year survival rate.

Hepatocyte Growth Factor Receptor (C-MET) [74-77]

[0159] C-MET is a proto-oncogene that encodes a transmembrane receptor tyrosine kinase for HGF. HGF is a mitogen for hepatocytes and endothelial cells, and exerts pleitrophic activity on several cell types of epithelial origin. The cellular localization of C-MET is the cell surface.

[0160] Hepatocyte growth factor/scatter factor (HGF/SF) stimulates a broad spectrum of epithelial cells causing them to proliferate, migrate, and carry out complex differentiation programs including angiogenesis. HGF/SF binds to a receptor encoded by the c-MET oncogene. While both normal and malignant tissues express the HGF receptor, expression of HGF/SF appears to be limited to malignant tissue.

[0161] While the human lung generally expresses low levels of HGF/SF, expression increases markedly in NSCLC. Using Western blot analysis, 88.5% of lung cancers exhibited an increase in the protein expression. All histologic types of tumors expressed the protein at increased concentrations. While increased levels of protein occur in all stages of the disease, recent evidence suggests that in addition to the cancer cells, stromal cells and/or inflammatory cells may be responsible for the production of the growth factor.

Mucin - MUC-1 [78-82]

[0162] Mucin-1 comes from a family of highly glycosylated secretory proteins which comprise the major protein constituents of the mucous gel which coats and protects the tracheobronchial tree, gastrointestinal tract and genitourinary tract. Mucin-1 is atypically expressed in epithelial tumors. The cellular localization of Mucin-1 is cytoplasm and the cell surface.

[0163] Mucins are a family of high molecular weight glycoproteins that are synthesized by a variety of secretory epithelial cells that are either membrane bound or secreted. Within the respiratory tract, these proteins contribute to the mucus gel that coats and protects that tracheobronchial tree. Changes in mucin expression commonly occur in conjunction with malignant transformation including lung cancer. Evidence exists suggesting at these changes may contribute to alterations in cell growth regulation, recognition by the immune system, and the metastatic potential of the tumor.

[0164] Although normal lung tissue expresses MUC-1, significantly higher levels of expression are found in lung cancer with highest levels occurring in adenocarcinoma. Staining appears to occur independently of stage and is more common in smokers than in former smokers or nonsmokers. Some premalignant lesions also exhibit increased MUC-1 expression.

Thyroid Transcription Factor-1 (TTF-1) [83,84]

[0165] TTF-1 belongs to a family of homeodomain transcription factors that activate thyroid-specific and pulmonary-specific differentiation genes. The cellular localization of TTF-1 is nuclear.

[0166] TTF-1 is a protein originally found to mediate the transcription of thyroglobulin. Recently, TTF-1 expression was also found in the diencephalon and brohchioloalveolar epithelium. Within the lung TTF-1 functions as a transcription factor regulating the synthesis of surfactant proteins and clara secretory protein. Overexpression of TTF-1 occurs in a large proportion of lung adenocarcinomas and can aid in distinguishing between primary lung cancer and cancers that metastasize to the lung. Adenocarcinomas that express TTF-1 and are cytokeratin 7 positive and cytokeratin 20 negative can be detected with 95% sensitivity.

Vascular Endothelial Growth Factor (VEGF) [33,61,85-89]

[0167] VEGF plays an important role in angiogenesis, which promotes tumor progression and metastasis. There are multiple forms of VEGF; the two smaller isoforms are secreted proteins and act as diffusible agents, whereas the larger two remain cell associated. The cellular localization of VEGF is cytoplasmic, cell surface, and extracellular matrix.

[0168] Vascular Endothelial Growth Factor (VEGF) is an important angiogenesis factor and endothelial cell-specific mitogen. Angiogenesis is an important process in the latter stages of carcinogenesis, tumor progression and is particularly important in the development of distant metastasis. VEGF binds to a specific receptor Flt that is often present in the tumors expressing the growth factor suggesting the presence of an autocrine loop.

[0169] Immunohistochemical analysis reveals that cells expressing VEGF exhibit a pattern of staining that is diffuse and cytoplasmic. While not expressed by nonneoplastic cells, VEGF is present in the majority of NSCLC and in a smaller percentage of SCLC. Several reports have shown high levels of VEGF in early stage lung cancer.

[0170] Expression of VEGF has been associated with an increased frequency of metastasis. Studies have shown that VEGF expression is indicative of a poor prognosis and shorter disease-free interval in adenocarcinoma but not in squamous cell carcinoma. Three year and five year survival rates in the group expressing high levels of VEGF were 50% and 16.7% as compared to 90.9 and 77.9% respectively for the low VEGF group.

Epidermal Growth Factor Receptor (EGFR) [90-104]

[0171] Epidermal Growth Factor Receptor (EGFR) is a transmembrane glycoprotein, which can bind and become activated by various ligands. Binding initiates a chain of events that result in DNA synthesis, cell proliferation, and cell differentiation. EGFR has been demonstrated in a broad spectrum of normal tissues, and EGFR overexpression is found in a variety of neoplasms. Increased expression has been observed in adenocarcinomas of the lung and large cell carcinomas but not in small cell lung carcinomas. The cellular localization of EGFR is the cell surface.

[0172] The EGFR plays an important role in cell growth and differentiation. The EGFR is uniformly present in the basal cell layer but not in more the superficial layers of histologically normal bronchial epithelium. With this exception, there is no consistent staining of normal tissue. Recent evidence suggests that the overexpression of the EGF receptor may not be an

absolute requirement for the development of invasive lung cancer. However, it appear that in cases where EGFR overexpression occurs it is a relatively early event with greater staining intensity in more advanced disease.

[0173] For patients with invasive carcinomas, 50-77% of tumors stain for EGF.

Overexpression of the EGFR is more common in squamous cell carcinoma than in adenocarcinoma and common in SCLC. Highest levels of EGFR occur in conjunction with late stage and metastatic disease that have approximately twice the concentration of EGFR as that seen in stage I/II tumors. Estimates suggest that the level of the EGFR observed in stage I tumors is approximately twice that seen in normal tissue. In addition, 48% of bronchial lesions also show EGFR staining including, metaplasia, atypia, dysplasia, and CIS. In the "normal" bronchial mucosa, of these same cancer patients, overexpression of the EGFR was observed in 39% of cases but was absent in the bronchial epithelium of the non-cancer. In addition, overexpression of the EGFR occurs more frequently in the tumors of smokers than in nonsmokers, particularly in the case of squamous cell carcinoma.

[0174] While several studies have suggested that overexpression of the EGFR is associated with the poor prognosis, other studies have failed to make this correlation.

Nucleoside Diphosphate Kinase/nm23 [105-111]

[0175] Nucleoside diphosphate kinase (NDP kinase)/nm23 is a nucleoside diphosphate kinase. Tumor cells with high metastatic potential often lack or express only a low amount of nm23 protein, hence the nm23 protein has been described as a metastasis suppressor protein. The cellular localization of nm23 is nuclear and cytoplasmic.

[0176] Expression of nm23/nucleoside diphosphate/kinase A (nm23) is a marker of tumor progression where there is an inverse relationship between expression and metastatic potential. In cases where stage I tumors overexpress nm23, no evidence of metastasis was seen during an average follow-up period of 35 months. Immunohistochemical analysis reveals staining that is diffuse, cytoplasmic and generally limited to malignant cells. Alveolar macrophages also express the protein. Given that high levels of expression are associated with a low metastatic potential, there is currently no explanation as to why normal epithelial cells do not express nm23.

[0177] Intense staining has been observed in high percentage of NSCLC particularly large cell lung cancer and 74% of SCLC suggesting that this protein plays an important role in tumor progression. With the exception of squamous cell carcinoma, staining intensity tends to increase

with stage. Based on the available evidence, it would appear that nm23 is a prognostic factor in both SCLC and NSCLC.

Bcl-2 [101,112-125]

[0178] Bcl-2 is a mitochondrial membrane protein that plays a central role in the inhibition of apoptosis. Overexpression of bcl-2 is a common feature of cells in which programmed cell death has been arrested. The cellular localization of Bcl-2 is the cell surface.

[0179] Bcl-2 is a protooncogene believed to play a role in promoting the terminal differentiation of cells, prolonging the survival of non-cycling cells and blocking apoptosis in cycling cells. Bcl-2 can exist as a homodimers or can form a heterodimer with Bax. As a homodimer, Bax functions to induce apoptosis. However, the formation of a Bax-bcl-2 complex blocks apoptosis. By blocking apoptosis, bcl-2 expression appears to confer a survival advantage upon affected cells. Bcl-2 expression may also play a role in the development of drug resistance. The expression of bcl-2 is negatively regulated by p53.

[0180] Immunohistochemistry analysis of bcl-2 reveals a heterogeneous pattern of cytoplasmic staining. In adenocarcinoma, expression of bcl-2 was significantly associated with smaller tumors (<2 cm) and lower proliferative activity. The expression of bcl-2 appears to be more closely associated with neuroendocrine differentiation and occurs in a large percentage of SCLC. [0181] Overexpression of bcl-2 is not present in preneoplastic lesions suggesting that changes in bcl-2 occur relatively late in tumor progression. In addition to tumor cells, bcl-2 immunostaining also occurs in basal cells and on the luminal surfaces of normal bronchioles but is generally not detected in more differentiated cell types.

[0182] Association of bcl-2 immunoreactivity with improved prognosis in NSCLC is controversial. Several reports of suggested that patients with tumors expressing bcl-2 have a superior prognosis and a longer time to recurrence. Several reports indicate that bcl-2 expression tends to be lower in those patients who develop metastatic disease. For patients with squamous cell carcinoma, expression of bcl-2 has been linked to an improvement in 5-year survival. However, in three relatively large studies there was no survival benefit linked to bcl-2 expression, particularly for patients with early stage disease.

Estrogen Receptor-related Protein (p29) [126]

[0183] ER related protein p29 is an estrogen-related heat shock protein that has been found to correlate with the expression of estrogen-receptor. The cellular localization of p29 is cytoplasmic.

[0184] Estrogen-dependent intracellular processes are important in the growth regulation of normal tissue and may play a role in the regulation of malignancies. In one study expression of p29 was detected in 109 (98%) of 111 lung cancers. The relation between p29 expression and survival time was different for men and women. Expression of p29 was associated with poorer survival particularly in women with Stage I and II disease. There was no correlation between p29 expression and long-term survival in men.

Retinoblastoma Gene Product (Rb) [68,73,123,127-141]

[0185] Retinoblastoma Gene Product (Rb) is a nuclear DNA-binding phosphoprotein. Under phosphorylated Rb binds oncoproteins of DNA tumor viruses and gene regulatory proteins thus inhibiting DNA replication. Rb protein may act by regulating transcription; loss of Rb function leads to uncontrolled cell growth. The cellular localization of Rb is nuclear.

[0186] Retinoblastoma protein (pRb) is a protein that is encoded by the retinoblastoma gene and is phosphorylated and dephosphorylated in a cell cycle dependent manner. pRb is considered an important tumor suppressor gene that functions to regulate the cell cycle at G0/G1. In its hypophosphorylated state, pRb inhibits the transition from G1 to S. During G1, inactivation of the growth suppressive properties of pRb occurs when the cyclin dependent kinases (CDK's) phosphorylate the protein. The hyperphosphorylation of pRb prevents it from forming a complex with E2F that functions as a transcription factor proteins that are required for DNA synthesis.

[0187] Inactivation of the retinoblastoma (Rb) gene has been documented in various types of cancer, including lung cancer. Small-cell carcinomas fail to stain for pRb indicating loss of Rb function. Overall, 17.6% of the tumors fail to express pRb with no correlation being seen with respect to stage or nodal status. A reduction in staining has also seen in 31% dysplastic bronchial biopsies. However, there appears to be no correlation between pRb expression and the severity of dysplasia. In contrast, normal bronchial epithelium and cells taken from areas adjacent to tumors expressed pRb positive nuclei. These data suggest that alterations in the expression of the Rb protein may arise early in the development of some lung cancers.

[0188] Patients with Rb-positive carcinomas tend to have a somewhat better prognosis but, in most studies, the difference is not significant. However, patients with adenocarcinoma whose tumors are both pRb negative and either p53 or ras positive exhibit a decrease in 5-year survival. A similar relationship does not occur in squamous cell carcinoma. pRb negative tumors have been reported to be more likely to exhibit resistant to doxorubicin than Rb-positive carcinomas.

Thrombomodulin [142-147]

[0189] Thrombomodulin is a transmembrane glycoprotein. Through its accelerated activation of protein C (which in turn acts as an anticoagulant by binding protein S and thrombin), synthesis of TM is one of several mechanisms important in reducing clot formation on the surface of endothelial cells. The cellular localization of thrombomodulin is the cell surface.

[0190] Aggregation of host platelets by circulating tumor cells appears to play an important role in the metastatic process. Thrombomodulin plays an important role in the activation of the anticoagulant protein C by thrombin and is an important modulator of intravascular coagulation. In addition to its expression in normal squamous epithelium, expression of thrombomodulin also occurs in squamous metaplasia, carcinoma in situ, and invasive squamous cell carcinomas. Although present in 74% of primary squamous cell carcinomas, only 44% of metastatic lesions stained for thrombomodulin. These data suggest that, with progression, there is a decrease in thrombomodulin expression. Higher levels of expression tend to occur in well and moderately differentiated tumors when compared to poorly differentiated tumors.

[0191] Patients with thrombomodulin-negative squamous cell carcinoma tend to have a worse prognosis. Eighteen percent of patients with thrombomodulin-negative have a five-year survival as compared to 60% in cases where the tumors stained positive for the protein. Progression to metastatic disease was also more common in thrombomodulin-negative tumors (69% vs. 37%) and there was a greater tendency for these tumors to develop at extrathorasic sites. Thus, loss of thrombomodulin expression appears to be prognostic in cases of squamous cell carcinoma. The observation that changes in thrombomodulin expression occur in later stages of NSCLC and that the protein is expressed by normal bronchial epithelial cells would tend to limit its utility as a marker for early detection. However, since a majority of mesotheliomas and only a small percentage of adenocarcinomas express thrombomodulin, the marker has potential utility in discriminating between these two tumor types.

E-cadherin & N-cadherin [148-151]

[0192] E-cadherin is a transmembrane Ca2+ dependent cell adhesion molecule. It plays an important role in the growth and development of cells via the mechanisms of control of tissue architecture and the maintenance of tissue integrity. E-cadherin contributes to intercellular adhesion of epithelial cells, the establishment of epithelial polarization, glandular differentiation, and stratification. Down-regulation of E-cadherin expression has been observed in a number of carcinomas and is usually associated with advanced stage and progression. The cellular localization of E-cadherin is the cell surface.

[0193] E-cadherin is a calcium-dependent epithelial cell adhesion molecule. A decrease in E-cadherin expression has been associated with tumor dedifferentiation and metastasis and decreased survival. Reduced expression has been observed in moderately and poorly differentiated squamous cell carcinoma and in SCLC. There was no change in E-cadherin expression in adenocarcinoma. Furthermore, while adenocarcinomas express E-cadherin theses tumors fail to express N-cadherin which is in contrast to mesotheliomas that express N-cadherin but not E-cadherin. Thus, these markers can be used to discriminate between adenocarcinoma and mesothelioma.

[0194] Expression of E-cadherin can also be used to assess the prognosis of patients with squamous cell carcinoma. Whereas 60% of patients with tumors expressing E-cadherin survived three-year survival, only 36% of patients exhibiting a reduction in expression survived 3 years.

MAGE-1 and MAGE-3 [152-156]

[0195] Melanoma Antigen-1 (MAGE-1) and Melanoma Antigen-3 (MAGE-3) are members of a family of genes that are normally silent in normal tissues but when expressed in malignant neoplasms are recognized by autologous, tumor-directed and specific cytotoxic T cells (CTL's). The cellular localization of MAGE-1 and MAGE-3 is cytoplasmic.

[0196] MAGE-1, MAGE-3 and MAGE 4 gene products are tumor-associated antigens that are recognized by cytotoxic T lymphocytes. As such, they could have utility as targets for immunotherapy in NSCLC. MAGE proteins are also expressed by some SCLCs but not by normal cells. While the frequency of MAGE expression falls below the level necessary for use as a detection marker, differences in the pattern of expression between histologic types suggest that MAGE expression may have utility as differentiation markers. This utility is also supported by the observation that, in 50% of squamous cell carcinoma greater than 90% of tumor cells

showed evidence of MAGE-3 overexpression with 30% to tumors exhibiting overexpression in at least 50% of cells.

Nucleolar Protein (p120) [157]

[0197] p120 (proliferation-associated nucleolar antigen) is found in the cells of nucleoli of rapidly proliferating cells during early G1 phase. The cellular localization of p120 is nuclear.

[0198] Nucleolar protein p120 is a proliferation-associated protein whose function has yet to be elucidated. Strong staining has been detected in tumor tissue but not in macrophages or normal tissue. Overexpression of p120 was more common in squamous cell carcinoma that in adenocarcinoma or large cell carcinoma raising the possibility that this marker may have utility in discriminating between tumor types.

Pulmonary Surfactants [83,158-166]

[0199] Pulmonary surfactants are a phospholipid-rich mixture that functions to reduce the surface tension at the alveolar-liquid interface, thus providing the alveolar stability necessary for ventilation. Surfactant proteins appear to be expressed exclusively in the airway and are produced by alveolar type II cells. In the non-neoplastic lung, pro-surfactant-B immunoreactivity is detected in normal and hyperplastic alveolar type II cells and some non-ciliated bronchiolar epithelial cells. Sixty percent of adenocarcinomas contained strong cytoplasmic immunoreactivity with 10-50% of tumor cells exhibiting staining the majority of cases. Squamous cell carcinoma and large cell carcinoma failed to stain for pro-surfactant-B. [0200] Surfactant Apoprotein B (SP-B) is one in four hydrophobic proteins that make up the pulmonary surfactant, which is a phospholipid and protein complex secreted by type II alveolar cells. Squamous cell and large cell carcinomas of the lung and nonpulmonary adenocarcinomas do not express SP-B. The cellular localization of SP-B is cytoplasmic.

[0201] SP-A is a pulmonary surfactant protein that plays an essential role in keeping alveoli from collapsing at the end of expiration. SP-A is a unique differentiation marker of pulmonary alveolar epithelial cells (type II pneumocytes); the antigen is preserved even in the neoplastic state. The cellular localization of SP-A is cytoplasmic.

[0202] Pulmonary surfactant A appears to be specific for non-mucinous bronchoiolo-alveolar carcinoma with 100% staining as compared to none of the of mucinous type. Pulmonary surfactants potentially have utility in discriminating lung cancer from other cancers metastasized to lung. In addition to tumor cells, non-neoplastic pheumocytes also stain for pulmonary

surfactant A. As with pulmonary surfactant B staining for pulmonary surfactant A is relatively common in adenocarcinoma but not in other forms of NSCLC or in SCLC. Mesothelioma also fails to express pulmonary surfactant A leading to the suggestion that pulmonary surfactant A may have utility in the discrimination between adenocarcinoma and mesothelioma.

Ki-67

[0203] Ki-67 is a nuclear protein that is expressed in proliferating normal and neoplastic cells and is down-regulated in quiescent cells. It is present in G1, S, G2, and M phases of the cell cycle, but is absent in G0 phase. Commonly used as a marker of proliferation. The cellular localization of Ki-67 is nuclear.

Table 5: .

Marker	Squamous Cell		Large Cell	Small Ceil	
	Carcinoma	Adenocarcinoma	Carcinoma	Carcinoma	Mesothelioma
Gluti	100.0*	· 64.5	- 80.5	64.0	NDA*
Glut3	17.5	16.0	39.5	9.0	NDA*
HERA	100.0	· 100.0	100.0	NDA	4.5
Basic FGF	83.0	48.7	50.0	100.0	NDA
Telomerase	82.3	86.3	93.0	66.7	NDA
PCNA	80.0	69.8	87.7	51.0	NDA
CD44v6	79.3	34.8	44.2	0.0	NDA
Cyclin A	79.0	68.0	83.5	97.0	NDA
Cyclin D1	42.7	36.0	62.0	90.0	NDA
Hepatocyte Growth	75.5	78.3	100.0	NDA	100.0
Factor /Scatter Factor	, ,				
MUC-1	55.5	90.0	100.0 ·	100	NDA
TTF-1	38.0	76.0	NDA	83.0	NDA
VEGF	61.8	68.3	100.0	43.5	NDA
EGF Receptor	63.1	45.3	96.0	Frequently	NDA
nm23	68.0	52.6	83.5	73.5	NDA.
Bcl-2	45.5	43.3	42.5	92.0	NDA
Loss of pRb	20.1	25.8	35.4	85.3	NDA
Expression					l <u></u>
Thrombomodulin	66.8	12.2	4.0	0.0	81.0
E-cadherin	69.0	85.0	NDA	100.0	0.0
N-cadherin	. NDA	4.0	NDA	NDA	94.0
MAGE 1	45.0	35.0	NDA	16.5	NDA
MAGE 3	72.0	33.3	NDA	33.5	NDA
MAGE 4	45.5	11.0	NDA	50.0	NDA
Nucleolar Protein	68.0	35.0	30.0	NDA:	NDA
(p120)				**	
Pulmonary Surfactant	0.0	61.5	0.0	NDA	NDA
В			<u> </u>		
Pulmonary Surfactant	12.0	52.9	17.5	20	0.0
Α					J

^{*} percent of tumors exhibiting a change in marker expression

a. Obtaining a Library of Markers of a Suitable Size

[0204] Preliminary pruning steps were required in order to obtain a suitable size library of markers that were correlated with lung cancer. More than a hundred markers correlated to lung cancer are known in the literature. A partial listing of candidate probes identified in the literature and evaluated for potential inclusion in panels tests include antibodies to: bax, Bcl-2, c-MET (HGFr), CD44v4, CD44v4, CD44v5, CD44v6, cdk2 kinase, CEA (carcino-embryonic antigen),

^{*} No Data Available

Cyclin A, Cyclin D1, E-cadherin, EGFR, ER-related (p29), erbB-1, erbB-2, FGF-2 (bFGF), FOS, Glut-1, Glut-2, Glut-3, Glut-4, Glut-5, HERA (MOC-31), HPV-16, HPV-18, HPV-31, HPV-33, HPV-51, integrin VLA2, integrin VLA3, integrin VLA6, JUN, keratin, keratin 7, keratin 8, keratin 10, keratin 13, keratin 14, keratin 16, keratin 17, keratin 18, keratin 19, A-type lamins (A; C), B-type lamins (B1; B2), MAGE-1, MAGE-3, MAGE-4, melanoma-associated antigen clone NKI/C3, mdm2, mib-1 (Ki-67), mucin 1 (MUC-1), mucin 2 (MUC-2), mucin 3 (MUC-3), mucin 4 (MUC-4), MYC, N-cadherin, NCAM (neural cell adhesion molecule), nm23, p16, p21, p27, p53, p120, P-cadherin, PCNA, Retinoblastoma, SP-A, SP-B, Telomerase, Thrombomodulin, Thyroid Transcription Factor 1, VEGF, vimentin, and waf1. The initial list of markers was pruned by initially assessing, from the literature, the apparent effectiveness of the probes in detecting early stage cancer cells, discriminating between cells of differing cancer states, and localizing the label to the target cancer cells. This list of markers was further pruned by removing markers whose utilization would be difficult to reduce to practice because they are difficult to produce or obtain, have unsuitable detection technology requirements or poor reproducibility of reported results. After all of the pruning steps were complete, a library of 27 markers was obtained.

b. Optimizing Protocols and Obtaining Gold Standard Lung Cancer Samples [0205] Preliminary preparation steps were also required prior to obtaining the panels. The probes containing appropriate labels were available from commercial vendors. The protocols of the probes were analyzed for optimum objective quantitative detection. For example, it was determined that the concentration of PCNA was too low. Originally, PCNA was diluted 1:4000 in S809 buffer. A second dilution was made, which was 1:3200 in S809. The optimized protocols for each marker is shown in below. It is noted that the second column is labeled "Antibody Name". Except for MOC-31, the probes in this list are listed by the marker name because many of the vendors refer to the antibody by the name of the marker. It is noted that an alternative way these reagents might be listed is, for example, anti-VEGF, anti-Thrombomodulin, anti-CD44v6, etc.

[0206] Gold standard tissue specimens were obtained from UCLA. Tissue specimens were received from two sources. Cases had been diagnosed using standard procedures including review of hematoxylin and eosin (H&E)-stained slides and the clinical history. Specimen slides

were coded and labeled with arbitrary numbers to blind the study pathologists to the historical diagnosis and antibody marker and to protect patient confidentiality.

[0207] Specimen slides with tissue sections from cancerous and non cancerous (control) tissues were used. A total of 175 separate cases were analyzed. Within this set, the following diagnoses, located in Table 6 were present with the following frequencies:

Table 6:

	Diagnosis	Number of occurrences
	Adenocarcinoma	25
i i	Large Cell Carcinoma	18
Cancer	Mesothelioma	26
ບຶ	Small Cell Lung Cancer	20
	Squamous Cell Carcinoma	24
	Emphysema	34
Control	Granulomatous Disease	3
Co	Interstitial Lung Disease	25

c. Determination of the Level of Expression of the Panel of Molecular Markers [0208] Sufficient specimen slides were prepared for each case so that only one probe was tested per slide. In general, a microscope slide is prepared which contains the cytologic sample contacted with one or more labeled probes that are directed at particular molecular markers. Independently, each study pathologists examined an H&E-stained slide to make a diagnosis for each case, and then examined each probe-reacted and immunochemically-stained slide to assess the level of probe binding, recording the results on a standardized data form.

[0209] In greater detail, the immunohistochemical staining was performed on formalin fixed, paraffin embedded (FFPE) tissue. Tissue sections were cut at 4 microns thick on poly-L-Lysine coated slides and dried at room temperature overnight. De-paraffinization and rehydration of the tissue sections were performed as follows: To completely remove all of the embedding medium from the specimen the slides were incubated in two consecutive Xylene-substitute (Histoclear) baths for five minutes each. All liquid was tapped off the slides before incubation in two consecutive baths of 100% reagent grade alcohol for three minutes each. Once again all excess liquid was tapped off the slides before being incubated in two final baths of 95% reagent grade alcohol for three minutes each. After the last bath of 95% the slides were rinsed in tap water and

held in wash buffer (Tris-buffered saline wash buffer containing 0.05% Tween 20 corresponding to a 1:10 dilution of DAKO Autostainer Wash buffer, code S3306). Table 7, below, presents a complete list of the reagents used in this study along with corresponding product code numbers. Detection systems used in the study were DAKO EnVision+ HRP mouse (code K4007) or rabbit (code K4003) and LSAB+ HRP (code K0690). The protocols for immunoassaying were followed according to the package inserts. The kits contained liquid two component DAB+ substrate chromogen (code K3468).

Table 7:

Reagents used in the Study	1 .
Reagents	Code #
National Diagnostics HistoClear	HS-200
Mallinckrodt Reagent Alchohol Absolute	7019-10
DAKO Antibody Diluent	S809
DAKO Background Reducing Antibody Diluent	S3022
DAKO Autostainer Buffer 10X	S3306
DAKO Target Retrieval Solution	S1700
DAKO Hi pH Target Retrieval Solution	S3307
DAKO Proteinase K	S3020
Rite Aid Hydrogen Peroxide 3%	None
DAKO Protein Block Serum Free	X0909
DAKO Goat Serum	X0501
DAKO Swine Serum	X0901
DAKO EnVision+ Mouse	K4007
DAKO EnVision+ Rabbit	K4003
DAKO LSAB+	K0690
DAKO DAB+	K3468
DAKO Hematoxylin	S3302
Dakomount Mounting Media	S3025
Instruments	Serial Numbers
DAKO Autostainers	3400-6613-03
	3400-6142R-03
Autostainer IHC Software Version V3.0.2	

[0210] Pretreatments were critical in optimizing these antibodies on lung tissue. For antibodies requiring enzyme digestion, DAKO Proteinase K (code S3020) was used for 5 minutes at room temperature. Antibodies requiring heat induced target retrieval received pretreatment using either DAKO Target Retrieval Solution (code S1700) or DAKO High pH Target Retrieval Solution (code S3307). Tissues were placed in a pre-heated Target Retrieval Solution and

incubated in a 95°C water bath for 20 or 40 minutes depending on the specific protocol. Tissue sections were then allowed to cool at room temperature for an additional 20 minutes.

[0211] After de-paraffinization, rehydration and tissue pretreatment, all specimens were incubated in a solution of 3% hydrogen peroxide to quench endogenous peroxidase activity. Blocking reagents were used specifically for the two antibodies FGF and Telomerase in order to minimize nonspecific background.

[0212] As shown in Table 8, below, tissue specimens were incubated for a specified length of time with 200 micro liters of the optimally diluted primary antibody. It is noted that the numbering of the markers/antibodies in Table 8 is consistent with the numbering of the antibody probes and markers throughout this document. Slides were then washed in DAKO 1X Autostainer Buffer (code S3306). Depending on the antibody, the correct detection system was applied. The steps and total incubation times for the DAKO EnVision+ HRP and LSAB+ HRP detection systems are shown in Table 9, below. The color reaction is developed using 3,3'-diaminobenzidine (DAB) resulting in a brown color precipitate at the site of the reaction.

Table 8:

minde	ies for Lung Panel	 	+		 	_		 	
	Antibody to	 			 -				
2	Marker:	Pretreatment	Block	Oilution	Primary Inc	Detection Sys	Clone	Vendor	- Code#
1	VEGF	HI pH TRS 20 min \$3307	Nane	1:15 in \$809	30 minutes	EnV-mouse	JH121	NeoMarkers	MS-350-P
Z	Thrombomodulin	None	Nane	1:100 in 5809	30 minutes	EnV+mousa	1009	DAKO	M0817
3	CD44v8	TRS 20 min \$1700 .	None	RTU	30 minutes	EnV+mouse	VFF-7	NeoMarkers	MS-1093-R
4	SP-A	None .	None	1:200 in S809	30 ininutes	EnV+mouse	PE10	DAKO	JM4501
5	Retinoblastoma	TRS 40 min \$1700	None	1:25 In S809			Rb1	DAKO	M7131
6	E-Cacherin	TRS 20 min \$1700	None	1:100 in \$809		EnV+mouse	NCH-38	IDAKO	M3612
7	Cyclin A	TRS 20 min S1700	None	1:25 in S809		EnV+mouse	626	Novocastra	NCL 117205
a	nm23	Hi pH TRS 20 min \$3307	None	1:50 in S809	30 minutes	EnV+ rabbit	Potycional	DAKO	A0098
9	Teloinerase	TRS 20 min \$1700	Prot Block X0909, 30 min	1:400 in SB09	Overnight	EnV+ rabbit	Potvctonal	Alpha Olagnostic	EST21-A
			w/5% goat serum X0501						-
10	KI-87	TRS 40 min S1700	None	1:200 In S809	30 minutes	EnV+mousa	IVAK-2	DAKO	M7240
11	Cyclin D1	HI OH TRS 20 min \$3307	None	1:200 in S3022	30 minutes		DCS-8	DAKO	M7155
12	PCNA Dilution 1	TRS 20 min S1700	None	1:4000 in \$809	30 minutes	EnV+mouse	PC10	OAKO ·	M0879
13	MAGE-1	Hi pH TRS 20 mm \$3307	None	1:250 in \$809			MA454	NeoMarkers	MS 1087
14	Mucin I	TRS 20 min S1700	None	1:40 in S809			VU4H5	Santa Cruz Blotech	Sc-7313
15	SP-8	TRS 20 min \$1700	None	1:100 in \$809			SP802	NeoMarkers	MS-1300-P
16	HERA	TRS 40 min S1700	None	1:50 in S809	30 minutes		MOC-31	DAKO.	M3525
17	FGF-2	None	Pmt Block X0909, 30 min	1:50 in S809	Overnight	EnV+mouse	bFM-2	Upstate Blotech	#05-11B
			w/5% swine serum X0901	1	1				
18	C-Met	Incomplete	None	Incomplate	Incomplete	EnV+mouse	8F11	Novocastro	118405
19	TTF-I	TRS 40 min \$1700	None	1:25 in S809	30 minutes	EnV+mouse	8G7G3/1	OAKO	M3575
20	Bcl-2	Hi gH TRS 20 min \$3307	None	1:75 In S809	30 minutes	EnV+mouse	124 .	DAKO	M0887
21	0120	TRS 20 min \$1700	None	1;10 in S809	30 minutes	EnV+mouse '	FB-2	Biogenex	MU198-UC
22	N-Catherin	TRS 40 min S1700	None	1:75 in S809	30 minutes	EnV+mouse	6G4 & 6G11	DAKO	NA
23	EGFR	Prot K 1:25 for 5 min	None	1:1500 in 5809			2-18C9	OAKO	K1492
24	Glut 1	TRS 40 min S1700	None	1:200 in \$800	30 minutes		Polycional	Santa Cruz Blotech	SC 1605
25	ER-related (p29)	TRS 40 min \$1700	None	1:200 in S809	30 minutes		G3.1	Blogenex	MU171-UC
26	Mage 3	TRS 40 min \$1700	None	1:20 in \$809			57B	G. Spagnoli	N/A
27	Glut 3	TRS 20 min \$1700	None	1:80 in S809	30 minutes		Polycional	Santa Cruz Blotech	SC 7581
28	IPCNA Dilution 2	TRS 20 min \$1700	None	1:3200 in \$809			PC10	DAKO	M0879

Table 9:

Detection S	ystems Used in the Study		Ī		
Steps					
	1 Deparatinization and rehydration				
	2 baths of Histoclear for 5mins each				
	2 baths of 100% alchohol for 3 mins each				
·.	2 baths of 95% alchohol for 3 mins each				
	Water Rinse				
	2 Pretreatments				
	TRS 40 or 20 mins				
	High pH TRS 20 mins				
	Proteinase K for 5 mins				
•	Water Rinse				
	3 Peroxidase block				
	Peroxide bath for 5 mins				
	Water Rinse			T	
	Buffer for 5 mins		-		
	Protein Block for 30 mlns after H2O2 Block				
	4 Primary Ab	•	1.		
'1	30 mins or Overnight at room temp				
	5 Detection System				
	EnV+ Systems	Labelled Polymer	OR	LSAB+ System	Secondary Reagent
	·	30 mins			15 mins Secondary Ab link
			<u> </u>		
· <u> </u>			<u> </u>		Tertiary Reagent
				<u> </u>	15 mins SA-HRP
	6 Chromogen		ļ.:	<u> </u>	
		Chromogen	1		Chromogen
		10 mins DAB+	1		5 minsDAB+

[0213] Following immunostaining all slides were incubated in DAKO Hematoxylin (code S3302) for 3 minutes and coverslipped using DAKOMount Mounting Media (S3025). All protocols were run on DAKO Autostainers (serial #'s 3400-6612-03 & 3400-6142R-03) using the IHC software version 3.0.2.

[0214] Immunostaining was viewed under a light microscope to determine that controls were correctly stained and tissues were intact. Slides were labeled, boxed and sent to designated pathologists for results interpretation. Trained pathologists identified the type of cancer or other lesion seen in the samples. Trained pathologists assessed the sensitivity to the marker probe by estimating the staining density and proportion of cells stained. These scores were entered in a data sheet for that patient. The pathologists were blinded to the original diagnosis and antibody marker used in the immunostaining. Each slide was read by at least two pathologists and results recorded on a data collection form. To provide additional integrity to the process, the method is

repeated with a second or third pathologist. The scores obtained can then be matched to identify data entry errors. The additional data also facilitates a better classifier design.

[0215] For each case, up to 27 slides were analyzed, each stained for a marker coded with numbers 1 through to 17, 19 through to 28. Staining for marker 18 (C-MET) could not be optimized and the marker/probe was therefore not used. Pathologist 1 scored slides from all 175 cases. Pathologist 2 scored slides from 99 of the cases. Pathologist 3 scored slides from 80 of the cases.

[0216] Table 10 below shows how many cases of each diagnosis each pathologist scored slides from:

Table 10:

7	Diagnosis	Pathologist 1	Pathologist 2	Pathologist 3
	Adenocarcinoma	25	12	14
75	Large Cell Carcinoma	18	9	.9
Cancer	Mesothelioma	- 26	14	. 8
ర	Small Cell Lung Cancer	20	12	6
	Squamous Cell Carcinoma	24	13	11
	Emphysema	. 34	23	13
tro	Granulomatous Disease	. 3	3 .	2
Control	Interstitial Lung Disease	25	13	17

[0217] For the purposes of some selected statistical analysis techniques, it was necessary to consider only those cases that had scores for all 27 slides present. Table 11 below shows how many cases of each diagnosis were complete in terms of having scores from all 27 slides.

Table 11:

	Diagnosis	Pathologist	Pathologist	Pathologist
	_	1	2 .	3
	Adenocarcinoma	14	10	8
15	Large Cell Carcinoma	12	9	3
Cancer	Mesothelioma	17	13	3
ပြီ	Small Cell Lung Cancer	7	9	1
	Squamous Cell Carcinoma	12	13	4 .
	Emphysema	32	21	1 .
tro	Granulomatous Disease	2	1	0 .
Control	Interstitial Lung Disease	23	7.	3

[0218] From this table, it can be calculated that each pathologist scored the following total number of complete cases. Pathologist 1 scored all 27 slides for 119 of the cases Pathologist 2 scored all 27 slides for 83 of the cases. Pathologist 3 scored all 27 slides for 23 of the cases. [0219] The total number of cancer data points is 172. This comprises 113 data points from Pathologist 1 and 60 data points from Pathologist 2. The total number of control data points is 101. This comprises 62 data points from Pathologist 1 and 39 data points from Pathologist 2. [0220] Figure 3 shows a comparisons between H-scores for probes 7 and 15 in control tissue and in cancerous tissue. The x-axis shows the H-scores while the y-axis shows the percent of cases with that particular H-score. The difference in H-scores is apparent. [0221] For each patient the scores were entered electronically into a Pathology Review Form which consolidates the scores into a data base showing the patient identifier together with diagnosis, proportion of cells stained, and staining density. The proportions and density were consolidated into a single "H-Score" obtained by grading the intensity as: none = 0, weak = 1, moderate = 2, intense = 3, and the percentage cells as: 0.5% = 0, 6.25% = 1, 26.50% = 2, 51-75% = 3, >75% = 4, and then multiplying the two grades together. For example, 50% weakly stained plus 50% moderate stained would score $10 = 2 \times 2 + 2 \times 3$. This is the standard scoring system throughout the analysis, except for the section 3(f), below, titled "Effect of Using other (non-H-score) objective scoring parameters", which investigates alternative scoring systems. [0222] Standard classification procedures were used to find the best combination of probes. Typically these use a search procedure such as the "Branch and Bound Algorithm" to find a hierarchy of the best features, ranked according to a test of discriminating power, and truncated according to a test of significance. This process also defines the decision rule or rules for best

[0223] The performance of a classifier designed with these features can be estimated from the data used to design the classifier. The straightforward application of all the design data to the classifier gives a very unsound estimate of performance.

classification.

[0224] The analysis of the data collected in the present example provide the optimum selection of probes which provided the best separation of classes. Therefore, panels were obtained that only needed a few probes to perform the analysis. However the data showed that near-optimum performance could be obtained with other combinations of probes. Hence, the invention is

flexible in being adaptable to the availability of probes where cost or supply problems may not allow the very best combination. In some cases, the invention can simply be applied to the available features to find an alternative combination. In other cases, the algorithm may be used to select features which allows cost weightings to be included in the selection process to arrive at a low cost solution.

[0225] The design of data collection and analysis experiment was chosen to avoid biases through the well established double blind procedures where data collection and data analysis were done independently.

[0226] In the first case the pathologists reviewed slides with conventional staining to allow a diagnosis to be made. This diagnosis was entered on the Pathology Review form. The pathologists were then presented, in random order, with slides stained by the marker probes for scoring the percentage of cells stained and the relative intensity of the staining. The slides were numbered to exclude information about the probe from the pathologist. To allow data integrity to be checked two pathologists reviewed all patients.

[0227] Data were consolidated into a database that was then reviewed by a team of statisticians. Probes were numbered to render their method of action as unseen during the analysis of their effectiveness.

[0228] The first stage of the analysis was to check the integrity of the data by comparing entries for each patient. Where large differences were found, the data entries were checked and any obvious errors were corrected. Unexplained differences were left in the data.

[0229] The data were then separately analyzed by four statisticians, using different techniques in recognition of the fact that different statistical methodologies are suited to different types of discriminating information in the data.

[0230] The first step in the process of selecting the best probe combination is to divide the data into two sets, one for designing a classifier and one for testing the performance of the classifier. By selecting the design made with the design (train) set, but showing the best performance evaluated on the test set, it can be concluded with confidence that the classifier has generalized to the structure of the data and not adapted to particular cases seen in the training set.

[0231] In order to test for reliability the analysis was typically repeated with many randomly selected sets of training data and test data. This approach is generally accepted as giving good

estimates of the classifier performance. Where these tests showed inconsistent selections of probes such probe selections were discounted as unreliable.

- d. Statistical Analysis and/or Pattern Recognition
- 1. Introduction to Data Analysis
 - a. Input Data
 - i. Raw Data
- [0232] For each patient the scores were entered electronically into a Pathology Review Form that consolidates the scores into a database showing the patient identifier together with diagnosis, proportion of cells stained, and staining density.
 - ii. Computed data
- [0233] The efficiency of the score for each probe used in the analysis is computed from the intensity/percentage tables. The proportions and density are consolidated into a single "H-Score" with a simple rule H= proportion stained x (3 if intense+2 if moderate+1 if weakly stained). This is the feature value associated with that probe.
 - iii. Alternative computed data parameters
- [0234] The H-score described above was heuristically derived, a simple analysis to find a better way of combining percentages and intensity failed to show a significant improvement over H-score (Section 3(f), titled "Effect of Using other (non-H-score) objective scoring parameters"). A larger data base may allow the extraction of a better rule in future.
 - iv. User supplied weighting criteria per marker
- [0235] The invention is flexible in being adaptable to the availability of features where cost or supply problems may not allow the very best combination. For example, the invention can simply be applied to the available features to find and alternative combination. Alternatively, the algorithm used to select features allows cost weightings to be included in the selection process to arrive at a minimum cost solution. Marker performance estimates are shown for combinations selected from all the markers collected or only those from one supplier. It is also shown how the C4.5 package can be used to down weight certain probes, say on the basis of their high cost. These probe combinations do not perform as well as the optimum combination, but the performance might be acceptable in circumstances where cost is a significant factor.

v. User supplied weighting criteria per class

[0236] Some of the methods used allow weightings to be applied to the classes. This is available in C4.5 where the tree design can optimize the cost. Also the Discriminant Function method gives a single parameter output which can be used to give a desired false positive or false negative probability. A plot of these parameters for different threshold settings is known as the Receiver Operating Curve.

vi. Detection panels – assumptions

[0237] A low probability of a false negatives was assumed to be desirable for the cancer detection process (to avoid positive patients being missed at the cost of an increased number of false positives who would require re-screening). It was also assumed that the cancer discrimination process would require a lower false positive score (to minimize patients receiving the wrong treatment).

[0238] It was assumed that detection panels requiring 6 or more probes to achieve an acceptable performance would not be cost effective. It was also assumed that a detection panel with a false negative error rate of more than 5% would not be acceptable. Panels falling outside this box are not accepted. This assumption acknowledges that cytometric panels are likely to have a worse performance than the histology based panels analyzed here. The ultimate aim will be a cytometric panel which performs better than 20% error rate, this being approximately the performance of cervical Pap smear screeners.

vii. Discrimination panels – assumptions

[0239] It was assumed that panels requiring 6 or more probes are not cost effective and it was assumed that an error rate of better than 20% is required. Panels falling outside this box were not accepted.

b. Output data

Outputs provided by the present analysis included:

Confusion Matrices, showing how data from the test set was classified as either true positive, false positive, true negative or false negative. These may be shown as actual counts or as percentages. Confusion matrices are discussed in section 2(d) titled "Performance Metrics". A confusion matrix shows how data from a test set was classified as either true positive, false positive, true negative or false negative. An exemplary confusion matrix, obtained from data analyzed by decision trees, is shown

below in table 12 for simultaneous discrimination of adenocarcinoma, squamous cell carcinoma, large cell carcinoma, mesothelioma and small cell carcinoma.

Table 12:

3	Adeno	Squamous	Large Cell	Mesolhelioma	Small Cell
Adeno	67.74%	6.45%	19.40%	0.00%	6.45%
Squamous Cell	2.94%	76.47%	11.67%	0.00%	8.82%
Large Cell	28.00%	8.00%	44.00%	8.00%	12,00%
Mesothelioma	0.00%	25.64%	51.28%	89.74%	2.56%
Small Cell	0.00%	3.85%	23.08%	3.85%	69.23%

- Error Rates, summarizing data in the confusion matrix as the sum of all false classifications divided by the total number of classifications made expressed as a percentage.
- Receiver Operating Characteristic (ROC) curves show the estimated percentage (or per unit probability) of false positive and false negative scores for different threshold levels in the classifier. An indifferent classifier, unable to discriminate better than random choice would present a ROC curve with equal true and false readings. The area under this curve would be 50% (0.5 probability).
- Area Under the Curve (AUC) is often used as an overall estimate of classifier
 performance and most standard discriminant function packages provide this AUC figure.
 A perfect classifier would have 100% Area Under the Curve, and a useless classifier
 would have an AUC near 50% (0.5 probability).
- Sensitivity and specificity (can be derived from the confusion matrix). See section 2(d)(iii) titled "Sensitivity and Specificity".
- Marker correlation matrices. See Figure 4.
 - i. Detection panels: composition

[0240] These panels are trained on data divided into two classes, patients with any of the five cancers and patients with none of the cancers. Not all probes were present for all patients. Where one or more probes were missing for a particular analysis these cases were excised from the data. Hence, where analysis was undertaken on reduced numbers of probes the data set might include slightly more cases.

[0241] The number of probes included in the analysis was 27. Although in many cases a false probe was added where the data entered for that probe was from a random number generator set

to generate numbers uniformly between zero and 12. This false probe was included in much of the early analysis to ensure integrity in the probe selection process. This false probe was also used in one approach to progressively eliminate probes from the analysis. Probes that contributed less information than the false probe could be readily identified and excluded from the selection process. Early elimination of such probes speeds the analysis and renders the analysis less vulnerable to variations in results (noise) caused by these probes.

ii. Detection Panel Performance

[0242] As outputs from this study, the probe combinations selected by the different methodologies and their performance estimates in terms of the confusion matrix, % error rate, and AUC are reported.

iii. Detection Panels – alternative compositions

[0243] Detection panels were also selected from reduced sets of probes. In one set of panels, performance measures of panels weighted for commercially preferred markers were obtained. The performances obtained when the best probe was removed from the analysis to find a new combination of discriminating probes was also analyzed. The performance of a single probe acting on its own was found to be very high (probe 7). However, as shown below in the performance diagrams, Table 13, evaluated using linear discriminant analysis, the performance was improved as more markers were added. The best subsets of probes were determined using best subsets logistic regression. The improvement is statistically significant.

Table 13:

Probe 7

4.	Cancer	Control
Cancer,	87.93%	12.07%
Control	0.00%	100.00%

Probes 7 and 16

語言語言	Cancer	Control
Cancer	93.10%	6.90%
Control	1.16%	98.84%

Probes 7, 15 and 16

To be the second	Cancer	Control
Cancer	90.52%	9.48%
Control	1.16%	98.84%

Probes 1, 7, 15, and 16

TO PARTY.	Cancer	Control
Cancer .	90.52%	9.48%
Control	0.00%	100.00%

Probes 1, 4, 7, 15, and 16

Teor State	Cancer	Control
Cancer	92.24%	7.76%
Control	1,16%	98.84%

[0244] The best and second best subsets of probes (determined using best subsets logistic regression) and evaluated using logistic regression is shown below. AUC=Area under ROC curve. It is noted that mean AUC is the average from 100 trials on random train and test partitions (70%:30%). The results are shown below, in Table 14.

Table 14:

Probes	Mean AUC
7	94.28
28	80.14
7, 16	95
7, 15	94.59
7, 15, 16	95.94
1, 7, 16	95.33
1, 7, 15, 16	95.61
4, 7, 15, 16	95.34
1, 4, 7, 15, 16	95.3
1, 7, 11, 15, 16	95.57

iv. Discrimination panels - composition

[0245] For this part of the study five classifiers were designed and tested, each designed to detect the presence of one of the cancer from all patients with cancer. The application of this five way pair-wise system allows doubtful cases to appear more than once in the analysis, or not at all. Such cases can be identified and subjected to closer scrutiny, re-testing or alternative testing regimes.

[0246] Again the number of probes in the study was 27, with a false probe used in the early stage to reduce the numbers in the analysis.

v. Discriminant panels – performance

[0247] The performance estimators described above were used to show the performance of the best probe combinations discovered by the different techniques.

vi. Discriminant panels – alternative composition

[0248] The analysis was repeated for a probe combination comprising commercially preferred probes. Performance was degraded, but not unusable for several reduced-set classifiers. Below, the best subsets of probes without probe 7, determined using best subsets logistic regression), is shown, as Table 15. The data was evaluated using linear discriminant analysis.

Table 15:

Probe 28

(新作号)建	Cancer	Control
Cancer	0.706897	0.293103
Control	0.093023	0.906977

Probes 10 and 28

	Cancer	Control
Cancer	0.793103	0.206897
Control	0.034884	0.965116

Probes 10, 15 and 28

建	Cancer	Control
Cancer	0.810345	0.189655
Control	0.011628	0.988372

Probes 1, 10, 15 and 28

	Cancer	Control		
Cancer	0.827586	0.172414		
Control	0.011628	0.988372		

Probes 1, 10, 15, 16 and 28

BOX PART	Cancer	Control
Cancer	0.827586	0.172414
Control	0.011628	0.988372

[0249] The best and second best subsets of probes with probe 7 (determined using best subsets logistic regression) and evaluated using logistic regression is shown below. AUC=Area under ROC curve. It is noted that mean AUC is the average from 100 trials on random train and test partitions (70%:30%). The results are shown below, in Table 16.

Table 16:

Probes	Mean AUC
28	79.36%
10	82.28%
10, 28	94.21%
15, 28	88.68%
· 10, 15, 28	92.90%
1, 10, 28	93.59%
1, 10, 15, 28	92.99%
8, 10, 15, 28	93.20%
1, 10, 15, 16, 28	93.13%
1, 8, 10, 15, 28	93.57%

2. Data analysis methodology

[0250] In this section, the process of gaining an initial understanding of the structure of the data as a guide to interpreting results from the different methodologies used is described.

a. Analysis of variance

Pathologist-to-pathologist variability and pooling pathologist

scores.

(1)
$$t$$
 - Test

[0251] Two pathologists reviewed each patient's slides in this clinical trial. Pathologist 1 reviewed all patients, Pathologist 2 also reviewed approximately half of this set and Pathologist 3 reviewed the remainder. With two independent estimates of the H-score, the consistency of pathologist performance could be tested.

[0252] A readily available statistical tool was used to test the variability between pathologists. This is the paired-sample t-test. This takes the difference between each pair of estimates, averages these and expresses this as a proportion of the overall variances. The t-test then converts this ratio into a probability estimating the likelihood that the two samples sets came from the same population (the P value).

[0253] This test was applied to the scores for each marker probe, for all cases reviewed by Pathologist 1 and Pathologist 2, and also for all cases reviewed by Pathologist 1 and Pathologist 3. Since there were 27 tests applied (to cover all probes) a low value of P=0.01 was selected as the "significant threshold". Results, showing the P scores for each probe, and for the two pairs of pathologists, are shown below, in Tables 17, 18,19 and 20. It is clear that Pathologist 1 and Pathologist 2 were more consistent than Pathologist 1 and Pathologist 3.

Table 17:

Pathologist 1, Pathologist 2 scores:

X!	X2	X3	X4	X5	X6	X7
0.5875446	0.01051847	0.4659704	0.4659704	0.3772394	0.2307273	0.01001357
	,					

X8 .	X9	X10 .	XII	X12	X13	·X14
0.004131056	0.7703014	0.1640003	0.2374452	0.9580652	0.1587876	0.001200265

X	15	X16	X17	X18	X19	X20 ·	X21
0.	19742	0.3860899	0.3829022	NA	0.544601	0.08873848	0.1686243

X22	X23	X24	X25	X26	X27	X28
0.5428451	0.1912477	0.4031977	0.2477236	0.5673386	0.9174037	0.00339071

Table 18:

Pathologist 1, Pathologist 2 scores thresholded at 0.01 ($\alpha = 1\%$ level of significance):

[]	Χī	X2	X3	X4.	X5	X6	X7
	TRU	TRU	TRUE	TRU	TRUE	TRU	TRUE
	E	E	*	E		E	

.X3	X9	X10	XII	X12	X13	X14
FAL.	TRU	TRU	TRU	TRU .	TRU	FALS
SE	ŀΕ	E	E	E	Е	E

ſ	X15	X16	X17	X13	X19	X20	X21
Ì	TR	TRU	TRU	NA	TRU	TRU	TRU
ļ	UE	E	E ·		E	Ε	Ε' .

ſ	X22	X23	X24	X25	X26	X27	X28
Ī	TR	TRU	TRU	TRU	TRU	TRU	FALS
-	UE	ĿΕ	E .	E	E	Ε	E

Table 19:

Pathologist 2, Pathologist 3 scores:

Xi .	X2	X3 ··	X4 .	X5	X6 .	X7
3.814506e-09	0.0399131	0.1954867	5.671062e-05	0.01856276 .	0.2757166	0.2292583
X8 .	X9	· X10	XII	X12	X13	X14
2.044038e-12	0.004166467	0.00983267	0.003710155	0.01461007	0.03312421	0.0003367823
•				× .		
X15	X16	X17	X18 .	X19	.X20	X21
0.0005162036	0.2276537	0.002987705		4.267708e-06	0.007287372	0.1654067
	,		•			
X22	X23	X24	X25	X26	X27 ·	X28 -
0.02400127	0.0009497766	2.478456e-07	0.1591684	0.08318303	3.122143e-05	

Table 20: Pathologist 1, Pathologist 3 scores thresholded at 0.01 ($\alpha = 1\%$ level of significance)::

ΧI	X2	X3	X4	X5	X6	1 X7
FALSE	TRUE	FALSE	FALSE	TRUE	TRUE	TRUE
			•	•		
X8	X9	X10	XII	-X12	X13	X14
FALSE	FALSE	FALSE	FALSE	TRUE	TRUE	FALSE
:	•					
X15	X16	X17	.X18	X19	X20	X21
FALSE	TRUE	FALSE	FALSE	FALSE	FALSE	TRUE
			٠.			
X22	. X23	X24	X25	X26	X27	X28
TRUE	FALSE	FALSE	TRUE	TRUE	FALSE	TRUE

[0254] Because the H score is subjective it is prone to scale factor differences and noise at marginal cases. So, in spite of the three features which showed statistically different scores between Pathologist 1 and Pathologist 2, this joint data was accepted as representative of a measuring instrument. Pathologist 1 and Pathologist 2 were combined into a single data set for the analysis process. The results for Pathologist 3 were withheld for independent testing purposes. Such tests using the Pathologist 3 data would be biased towards showing an underperformance because of the significant differences.

[0255] The data from Pathologist 1 and Pathologist 2 were combined by considering them as separate cases, with the variability giving a degree of independence between the results for any one case. When testing with such data the performance estimates will be biased towards a more optimistic value. This is because samples coming from the same patient may occur simultaneously in the training a test subsets. This does not however invalidate the processes used to find the best combination of features, it merely biases the estimate of performance.

(2) Analysis of Variance of H-Scores

(a) Background

[0256] Within each probe, the H-scores may vary due to many reasons. To the extent they vary consistently due to the type of disease this is useful, variation due to which pathologist read the slide is instructive, whereas random variation sets a limit on the detection of the previous two sources of variation.

[0257] Analysis of Variance (ANOVA) is a standard technique for splitting up the sources of variation in data and for testing its statistical significance. ANOVA summarizes the total

variation of a set of data as a sum of terms which can be attributed to specific sources, or causes, of variation.

[0258] ANOVA is available in many statistical packages. The public domain package "R" was chosen ("The R Project for Statistical Computing", http://www.R-project.org/).

(b) Aim

[0259] To perform ANOVA analyses on the H-score data from pathologists 1 and 2 and to consider whether this data can be safely merged into a single consistent set for further analysis for the selection of panels.

(c) Methodology

[0260] From the database, data was selected from pathologists 1 and 2. Only data which was complete for a given probe was used in the ANOVA for that probe.

[0261] The control categories of Emphysema, Granulomatous Disease, and Interstitial Lung Disease were grouped together and called "Normal" giving 6 levels within factor Disease.

[0262] Pathologist was coded as a factor with 2 levels (Pathologist 1, Pathologist 2).

[0263] An R script was written to perform a standard ANOVA analysis for each probe in turn, using the factors: Disease, Pathologist, and the interaction term Disease:Pathologist. The results are shown in below, in Table 21. "Df" is defined as the degrees of freedom. In a dataset of n observations, knowing n-1 deviations from the mean, the nth is automatically determined. N-1 is the number of degrees of freedom. Sum Sq and mean Sq are measures of variation. F is a test statistic concerning the equality of two variances based on the F distribution. Pr(>F) is the probability used to determine whether or not the variability is statistically significant.

Table 21:

Analysis of Variance of H-Scores

```
Probel

Df Sum Sq Mean Sq F value 2r(>F)

Disease 5 443.56 88.71 15.8202 3.690e-13 ***

Pathologist 1 0.66 0.66 0.1174 0.7323

Disease:Pathologist 5 15.34 3.07 0.5470 0.7405

Residuals 204 1143.93 5:61

---

Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1
```

```
Probe2

Df Sum Sq Mean Sq F value Pr(>F)

Disease 5 1067.39 213.48 24.1234 <2e-16 ***

Pathologist 1 13.02 13.02 1.4709 0.2263

Disease:Pathologist 5 27.98 5.60 0.6324 0.6752

Residuals 249 2203.50 8.85

---

Signif. codes: 0 `***' 0.001 `**' 0.05 `.' 0.1 ` ' 1
```

Prope3	
	Df Sum Sq Mean Sq F value Pr(>F)
Disease	5 1098.49 219.70 21.0751 <2e-16 ***
Pathologist	1 6.73 6.73 0.6458 0.4224
Disease:Pathologist	5 29.72 5.94 0.5703 0.7227
Residuals	243 2533.16 10.42
Signif. codes: 0 `***'	0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1

Probe4	
,	Df Sum Sq Mean Sq F value Pr(>F)
Disease	5 631.8 126.4 9.3707 3.454e-08 ***
Pathologist	1 6.6 6.6 0.4869 0.4860
Disease: Pathologist	5 13.1 2.6 0.1939 0.9647
Residuals	246 3317.1 13.5
Signif. codes: 0 '***'	0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1

```
Probe5

Df Sum Sq Mean Sq F value Pr(>F)

Disease 5 754.30 150.86 25.2826 <2e-16 ***

Pathologist 1 14.25 14.25 2.3875 0.1236

Disease:Pathologist 5 7.54 1.51 0.2528 0.9381

Residuals 248 1479.80 5.97

---

Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1
```

```
Probe6
                        Df Sum Sq Mean Sq F value
                                                      Pr(>E)
                            721.91 144.38 11.8515 2.771e-10 ***
Disease
                                     1.91 0.1568 0.6925
Pathologist .
                                      9.56 0.7850
                                                     0.5613
                         5
                             47.82
Disease: Pathologist
                       246 2996.93
                                     12.18
Residuals
Signif. codes: 0 '***' 0.001.'**' 0.01 '*' 0.05 '.' 0.1 ' 1
```

```
Probe7

Df Sum Sq Mean Sq F value Pr(>F)

Disease 5 1171.47 234.29 77.6802 < 2e-16 ***

Pathologist 1 8.84 8.84 2.9294 0.08847 .

Disease:Pathologist 5 46.36 9.27 3.0742 0.01063 *

Residuals 209 630.37 3.02

---

Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1
```

Probe8	· ·				1		
	Df	Sum Sq	Mean Sq	F value	Pr(>F)		
Disease	5	209.82	41.96	6.4352	1.201e-05	***	•
Pathologist	1	12.66	12.66	1.9407	0.16483		
Disease: Pathologist	· 5	71.20	14.24	2.1838	0.05654		
Residuals	251	1636.76	6.52		•		
		•					
Signif. codes: 0 `***'	0.0	01 `**' (0.01 '*'	0.05	' 0.1 ` '	L	

. Probe9			. •					
·					Pr(>F)			
Disease	. 5	197.21	39.44	8.4348	2.015e-07	***		
Pathologist	. 1	7.33	7.33	1.5681	0.2116			
Disease: Pathologist	5	24.56	4.91	1.0505	0.3884	•		
Residuals	265	1239.17	4.68	•				
				•				•
Signif codes: 0 `***'	0.0	01 `**'	0.01 `*'	0.05 .	0.1 ''	L .	 	

Probe10			
	Df Sum Sq Mean Sq F value Pr(>F)		
Disease	5 1113.46 222.69 39.0730 < 2e-16 ***		
Pathologist	1. 1.01 1.01 0.1778 0.67371		
Disease:Pathologist	5 62.45 12.49 2.1916 0.05635 .		
Residuals	213 1213.96 5.70		
	•	•	
Signif. codes: 0 `***'	0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1		*

```
Probell

Df Sum Sq Mean Sq F value Pr(>F)

Disease 5 320.15 64.03 9.5553 2.416e-08 ***

Pathologist 1 1.28 1.28 0.1918 0.6618

Disease:Pathologist 5 10.04 2.01 0.2996 0.9128

Residuals 245 1641.76 6.70

---

Signif. codes: 0 **** 0.001 *** 0.01 ** 0.05 ... 0.1 ' 1
```

```
Probel2

Df Sum Sq Mean Sq F value Pr(>F)

Disease 5 832.26 166.45 27.8793 <2e-16 ***

Pathologist 1 0.18 0.18 0.0307 0.8610

Disease:Pathologist 5 15.16 3.03 0.5079 0.7701

Residuals 248 1480.68 5.97

---

Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 `' 1
```

Probe13	Df	Sum Sa	Mean Sq	F value) Pr(>F)			
Disease		46.594	9.319	7.8408	8.674e-07	***	•	
Pathologist	1	0.044	0.044	0.0368	0.8481			
Disease: Pathologist	5	10.143	2.029	1.7069	0.1343			
Residuals	210	249.584	1.188					
Signif. codes: 0 '***	0.0	01 ***	0.01 `*'	0.05 .	0.1 '	1		- 0

```
Probel4

Df Sum Sq Mean Sq F value Pr(>F)

Disease 5 1305.69 261.14 23.9460 < 2e-16 ***

Pathologist 1 28.66 28.66 2.6279 0.10630

Disease:Pathologist 5 142.90 28.58 2.6208 0.02492 *

Residuals 243 2649.98 10.91

---

Signif. codes: 0 `***' 0.001 `**' 0.05 `.' 0.1 ` ' 1
```

```
Probel5

Df Sum Sq Mean Sq F value Pr(>F)

Disease 5 401.02 80.20 21.268 <2e-16 ***

Pathologist 1 13.17 13.17 3.493 0.0630 .

Disease:Pathologist 5 6.17 1.23 0.327 0.8963

Residuals 214 807.02 3.77

---

Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 `' 1
```

```
Probel6

Df Sum Sq Mean Sq F value Pr(>F)

Disease 5 2520.26 504.05 65.5572 <2e-16 ***

Pathologist 1 0.15 0.15 0.0194 0.8892

Disease:Pathologist 5 24.29 4.86 0.6318 0.6757

Residuals 247 1899.12 7.69

---

Signif. codes: 0 **** 0.001 *** 0.01 ** 0.05 ... 0.1 ' 1
```

```
Probel7

Df Sum Sq Mean Sq F value Pr(>F)

Disease 5 530.64 106.13 13.0178 2.426e-11 ***

Pathologist 1 8.42 8.42 1.0325 0.31050

Disease:Pathologist 5 109.96 21.99 2.6975 0.02131 *

Residuals 266 2168.55 8.15

---

Signif. codes: 0 **** 0.001 *** 0.01 ** 0.05 ... 0.1 ' 1
```

Probe19				• :			-	•	
	Df	Sum Sq	Mean Sq	F value	Pr (>F)				
Disease	5	1670.86	334.17	29.1960	<2e-16	***			
Pathologist	1	2.17	2:17	0.1895	0.6637	٠			
Disease:Pathologist	5	32.61	6.52	0.5698	0.7231			•	
Residuals	248	2838.56	11.45					•	
			•					,	
Signif. codes: 0 '***'	0.0	01 '**' (0.01 `*'	0.05 `.	0.1	' 1			

. Probe20		
	Df Sum Sq Mean Sq F value Pr(>F)	
Disease	5 964.71 192.94 34.2760 <2e-16 ***	·
Pathologist	1 8.83 8.83 1.5687 0.2116	
Disease: Pathologist	5 19.60 3.92 0.6963 0.6267	
Residuals	245 1379.12 5.63	
Signif. codes: 0 `***'	0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1	

```
Probe21

Df Sum Sq Mean Sq F value Pr(>F)

Disease 5 6.927 1.385 2.0604 0.07076.

Pathologist 1 0.464 0.464 0.6906 0.40670

Disease:Pathologist 5 1.576 0.315 0.4687 0.79945

Residuals 263 176.830 0.672

---

Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 `' 1
```

```
Probe22

Df Sum Sq Mean Sq F value Pr(>F)

Disease 5 640.16 128.03 31.7250 <2e-16 ***

Pathologist 1 1.64 1.64 0.4058 0.5247

Disease:Pathologist 5 18.78 3.76 0.9305 0.4617

Residuals 247 996.81 4.04

---

Signif. codes: 0 `***' 0.001 `**' 0.05 `.' 0.1 `' 1
```

Probe23			
		Sq F value Pr(>F)	
Disease		.12 46.5565 <2e-16	***
Pathologist	1 10.77 10.	.77 1.3092 0.2537	
Disease: Pathologist	5 20.92 4	.18 0.5084 0.7698	
Residuals 2	46 2024.39 8	. 23	
	•		
Signif. codes: 0 `***' 0	.001 '**' 0.01	`*' 0.05 `.' 0.1 `	' 1

Probe24				
	Df Sum Sq Mean Sq F value Pr(>F)			
Disease	5 516.06 103.21 24.0786 <2e-16 ***			
Pathologist	1 9.52 9.52 2.2210 0.1376			
Disease: Pathologist	5 12.48 2.50 0.5823 0.7135			·
Residuals	216 925.87 4.29	:		•
Signif. codes: 0 `***'	0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1		•	

Probe25	* *	
	Df Sum Sq Mean Sq F value Pr(>F)	
Disease	5 1761.26 352.25 34.5245 <2e-16 ***	
Pathologist	1 11.51 11.51 1.1285 0.2891	•
Disease: Pathologist	5 41.49 8.30 0.8134 0.5411	•
Residuals	248 2530.33 10.20	•
Signif. codes: 0 '***'	0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1	

Probe26		
	Df Sum Sg Mean Sg F value Pr(>F)	
Disease	5 399.85 79.97 13.6548 1.428e-11 ***	
Pathologist	1 0.30 0.30 0.0517 0.8204	
Disease:Pathologist	5 14.81 2.96 0.5056 0.7719	•
Residuals	214 1253.31 5.86	•
		.*
Signif. codes: 0 `***'	0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1	

```
Probe27
                         Df Sum Sq Mean Sq F value
                                                        Pr(>F)
                                              6.2551 1.956e-05 ***
                           5 117.92
                                      23.58
Disease
Pathologist
                               0.64
                                       0.64
                                              0.1695
                                                        0.6810
Disease: Pathologist
                           5
                              25.52
                                       5.10
                                             1.3539
                                                        0.2431
                         212 799.31
                                       3.77
Residuals
Signif. codes:
                0 '*** 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
Probe28
                             Sum Sq Mean Sq F value Pr(>F)
                          5 1634.60
                                     326.92 38.171 <2e-16 ***
Disease
                                       8.40
                                              0.981 0.3229
Pachologist
                               8.40
                            16.15
                                              0.377 0.8643
Disease:Pathologist
                                       3.23.
Residuals
                        267 2286.76
                0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1
Signif. codes:
```

(d) Analysis of Results

[0264] In all cases (except for probe 21) the response of the probes was related to disease. This is not surprising since the probes have presumably been selected for this purpose. In no case is the response of the probe related to pathologist (at the p=0.05 level). This indicates that it would be safe to merge this data and use the two pathologists as two measurements on the data.

[0265] In a few cases, probes 7, 14, 17, there is some evidence of an interaction term gaining significance. This indicates that there may be some difference between pathologists in their scoring of some diseases. Some of these cases may well be due to an occasional outlier in the data.

(e) Conclusions

[0266] The results indicate that it is safe to merge this data for further analysis. The data indicate that the slight interactions in some cases between pathologist and disease appear to be attributed to random sources.

ii. Patient to patient variability

[0267] The variability from patient to patient was measured by the disease: disease variability of section 2(a)(i)(2) (see above, "Analysis of Variance of H-Scores").

iii. Marker-to-marker variability

[0268] Histograms were plotted (PathologistData.xls, worksheet: Histograms) showing the distribution of marker scores for each probe for Control vs. Cancer.

b. Marker correlation matrix analyses

[0269] The population correlation coefficient ("Applied Mulitvariate Statistical Analysis", R. A. Johnson and D. W. Wichern, 2nd Ed,1988, Prentice-Hall, N.J.) measures the amount of linear association between a pair of random variables. Typically the distributions and associated parameters of the random variables are not known and the population correlation coefficient cannot be directly computed. In this case it is possible to compute the sample correlation coefficient from sample data. See Figure 4. The sample correlation coefficient is, however, only an estimate of the population correlation coefficient. Moreover, because it is calculated on the basis of sample data it is possible, purely by chance, that it may indicate a strong positive or negative correlation when in reality there may be no actual relationship between the corresponding random variables ("Modern Elementary Statistics", J. E. Freund, 6th Ed, 1984, Prentice-Hall, N.J.).

[0270] The correlation coefficient measures the ability of one variable to predict the other. A strong linear association does not, however, imply a causal relationship. The square of the correlation coefficient is called the coefficient of determination. The coefficient of determination computed for a bivariate data set measures the proportion of the variability in one variable that can be accounted for by its linear relationship to the other. When dealing with several variables, the correlation coefficient can be calculated for each pair in turn and the set of coefficients can be written as a matrix called the correlation matrix. See Figure 4.

[0271] The H-scores for the individual markers can be modeled as random variables. The sample correlation matrix for this multivariate data set can be computed from the input data described in the section titled "Input Data", above.

c. Pattern recognition

[0272] Statistical pattern recognition is an approach to classifying signals or geometric objects on the basis of quantitative measurements (called features). Statistical pattern recognition essentially reduces to the problem of dividing the *n*-dimensional feature space into regions that correspond to the categories or classes of interest.

[0273] Three different classifier methodologies employed in this study are sensitive to different structural forms within the data.

- [0274] For the Decision Tree method a preliminary analysis of different data combinations identified markers which were never used by C4.5 for the detection panel. These were removed from the analysis and this resulted in more consistent results, symptomatic of the left-out probes only contributing noise to the selection process.
- [0275] Similarly a preliminary analysis of probes used in the detection panels identified the noisy probes for removal prior to the detailed analysis.
- [0276] The Linear Discriminant Function method in SPSS has built-in stepwise processes for reducing the numbers of markers in the analysis. Typically, this reduced the probes used in the analysis to between 2 and 7.
- [0277] The Logistic Regression method in R and SAS implement stepwise procedures for variable selection. In SAS, a best subsets variable selection option is also provided. In R, the stepwise methodology was used in conjunction with multiple random trials to develop a heuristic method for selecting variables based on the number of times a given feature was used in 100 random selections of training and test data (split 70%:30% respectively). Features with counts comparable to the count for artificial random feature were progressively eliminated until a minimal consistent set of features was obtained over 100 runs.

i. Statistical methods

[0278] From the point of view of multivariate statistical analysis, the problem is one of estimating density functions in high-dimensional space (and partitioning this space into the regions of interest). Assuming that the distributions of random (feature) vectors are known, the theoretically best classifier is the Bayes classifier because it minimizes the probability of classification error (K. Fukunaga, "Statistical Pattern Recognition", 2nd Ed., Academic Press 1990, p.3). Unfortunately the implementation of the Bayes classifier is difficult because of its complexity, especially when the dimensionality of the feature space is high. In practice, simpler parametric classifiers are used. Parametric classifiers are based on assumptions about the underlying density or discriminant functions. The most common such classifiers are linear and quadratic classifiers. In multivariate statistical analysis such classifiers fall under the heading of discriminant analysis. Discriminant analysis techniques are closely related to multivariate linear

regression models and generalized linear models (encompassing, logistic and multinomial regression).

(1) Logistic Regression with a Binomial Response

(a) Background

[0279] The problem of selecting a set of markers to be used on a detection panel can be formulated as a logistic regression problem with a binomial response. The response variable is a factor with two levels: normal (no cancer) and abnormal (cancer). The explanatory variables are the marker H-scores.

[0280] The problem of selecting a set of markers to be used on a cancer discrimination panel can also be formulated as a logistic regression problem with a binomial response. The response variable is a factor with two levels: normal (not the cancer of interest) and abnormal (cancer of interest). The explanatory variables are the marker H-scores.

[0281] Stepwise variable selection can be used to select a subset of the original variables (markers) for use in discriminating between the two classes. This is a computationally expensive exercise and is best suited to a computer. Several commercial and public domain software packages---e.g., R, S-plus, and SAS---implement stepwise logistic regression.

[0282] Two different approaches to feature selection were investigated based on the stepwise variable selection procedures found in R and SAS respectively.

(b) Experimental data

[0283] The data used for the present analysis consists of the H-scores for markers 1-17, and 19-28 for the cases examined by Pathologist 1 and Pathologist 2 and described elsewhere in this report. In addition, a durnmy marker, 18, was added to the data set. The dummy marker consists of integer values from 0 to 12 selected at random from a uniform distribution.

(c) Method 1: Using the R package (version 1.4.1)

[0284] Computerized model fitting procedures generally cannot deal with missing data. This is the case for the glm (glm stands for generalized linear model) procedure used in R. Consequently when fitting a model using glm it was necessary to exclude all the cases for which there are one or more missing values. When fitting the initial full model, containing the 27 real markers and the single dummy marker, this reduces the data set to only 202 cases. With so few observations it was decided that the best way to perform variable selection, to train a classifier using the selected

variables, and to assess its performance was to undertake 100 trials on random partitions of the data into train and test sets.

- (i) Partitioning the data into train and test sets [0285] At the start of each trial, the data is partitioned into a test set and a training set. This is done by randomly choosing 30% of the abnormals and 30% of the normals to form the test set, and using the remaining observations to form the training set.
 - (ii) Variable (marker) selection

[0286] At the start of each trial, the full model, which includes all of the variables (markers), is fitted to the training data. In R the logistic regression model is fitted using glm. The code fragment used is as follows:

my.glm <- glm(my.model, family=binomial(link=logit)
data=training.data)</pre>

[0287] The procedure stepAIC is then used to perform stepwise variable selection based on the Akaike Information Criterion (AIC). This procedure is part of the publicly available MASS library. The library and the procedure are described in "Modern Applied Statistics with S-PLUS" (W. N. Venables and B.D. Ripley, Springer-Verlag, Pathologist 3 New York, 1999). The R code fragment to do this is as follows:

my.step <- stepAIC(my.glm, direction=both)</pre>

[0233] The resulting model is then assessed on the test data. The code fragment used is as follows:

```
probability_is_abnormal <-
predict(my.step,testing.data,type="response")</pre>
```

[0289] The performance of the classifier is recorded in terms of the actual error rate of misclassification (AER) and the area under the ROC curve (AUC). After the 100 trials, 100 models and their associated AERs and AUCs remain. A frequency table is constructed, recording the number of times each variable made an appearance in the 100 models. An example is shown in Table 22:

Table 22:

Variable	1	2	3	4	5	.6,	127.42	37	9	Til	12	14
Frequency	2	6	4	1	4	16	1.100	40	3	10	3.43.	1
Variable	15	16 73	17	18	119	20	22	23	24	, 25 3	28	
Frequency	4	,28-	2.	1	3	2	2	1-18	10	84	4	

[0290] This table is used to decide which markers to discard. First, all of the markers that have a frequency less than or equal to 10 are discarded. Next a cut-off frequency is chosen based on the frequency of the dummy marker (typically this is 1 or 1.5 times that of the dummy marker). All markers with a frequency less than this cut-off value are discarded. The remaining markers, along with the dummy marker, are then used as the full model for another 100 trials and the pruning process is repeated. If necessary, the severity of the pruning can be increased to force one or more markers out of the model. If necessary, the remaining markers can be used as the full model for yet another 100 trials. Pruning stops when the desired number of panel members is reached or the average AUC for the current model is less than that for the preceding model.

[0291] To illustrate the pruning process consider the table above. The table was obtained using the detection panel data. The shaded entries indicate those markers that are retained after pruning. Another 100 trials is performed using the following full model:

my.model <- Class $\sim X6 + X7 + X8 + X12 + X16 + X18 + X23 + X25$

[0292] Again, a frequency table, Table 23 is constructed:

Table 23:

Variable	# 6 m 17 7 7 18 18 112: 2 1 16	1 13	다. 23년 학생 1:25년 44
7 21 12010		'-	国际的 计控制线线
	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	17-	66 98
Frequency	1-63 - 1-1-100 - 1-5t - 1-48 : 1-30	47	66 98
	# 3 - 3 m (80)		* * * * * * * * * * * * * * * * * * * *

[0293] The shaded entries show the markers retained after pruning (using a cutoff of 47).

Another 100 trials is performed using the following full model:

my.mode1 <- Class $\sim X6 + X7 + X8 + X12 + X18 + X23 + X25$

[0294] Again, a frequency table, Table 24 is constructed:

Table 24:

Variable	6 7 . 7	8	12.5 / 18	23 - 25
Frequency	-96: 100	23	73: 3	88 98

[0295] At this point a cut-off of 50 is chosen. The shaded entries show the remaining markers for use on a 5 member panel. In each step, the average AUC increases: $94.37\% \rightarrow 95.45\% \rightarrow 95.78\%$.

(iii) Assessing the performance of the panel [0296] To assess the performance of the panel, 100 trials were performed, as before, but without the stepwise selection procedure. For each trial, the AUC, sensitivity, and specificity are recorded. For the detection panel example above, the results are:

> my.model <- Class $\sim X7 + X25 + X6 + X23 + X12$

> summary(AUC)

Min. 1st Qu. Median Mean 3rd Qu. Max. 0.9289 0.9590 0.9615 0.9601 0.9630 0.9630

> summary(sensitivity)

Min. 1st Qu. Median Mean 3rd Qu. Max. 0.8519 0.9630 0.9630 0.9737 1.0000 1.0000

> summary(specificity)

Min. 1st Qu. Median Mean 3rd Qu. Max. 0.8378 0.9730 0.9730 0.9749 1.0000 1.0000

[0297] In summary, the panel has a sensitivity of 97.37% and a specificity of 97.49%. The area under the ROC is 96.01%.

(d) Method 2: Using SAS (version 8.2)

[0298] Logistic regression can be performed in SAS using the procedure LOGISTIC. When the response variable is a two-level factor, the procedure fits a binary logit model (equivalent to glm in R with family=binomial and link=logit). SAS automatically excludes all of the missing multivariate observations for the model specified. Unlike R, SAS is able to perform a best subsets variable selection procedure. The code fragment in SAS needed to do this is as follows:

PROC LOGISTIC DATA=WORK.panel;

CLASS Class;

MODEL Class = X1 X2 X3 X4 X5 X6 X7 X8 X9 X10 X11 X12 X13 X14 X15 X16 X17 X18 X19 X20 X21 X22 X23 X24 X25 X26 X27 X28 /SELECTION= SCORE BEST=28; RUN;

[0299] This procedure is applied to the entire data set. The parameter BEST=28 directs SAS to find the best 28 single-variable models, the best 28 two-variable models, the best 28 three-variable models, up to the best 28 28-variable models.

(i) Assessing the performance of the panels [0300] The procedure described in method 1 is used to assess the performance of each of the panels. The following, Table 25, was generated from the detection panel data. It lists results only for the two best one-, two-, three-, four-, and five-marker panels.

Table 25:

Panel	Panel members	Sensitivity	Specificity	Area under ROC
1	7			94.28%
2	28			80.14%
3	7, 16	*		95.00%
4	7, 15			94.59%
-5	7, 15, 16			95.94%
6	1, 7, 16			95.33%
7	1, 7, 15, 16			95.61%
8	4, 7, 15, 16			95.34%
9	1, 4, 7, 15, 16			95.30%
10	1, 7, 11, 15, 16			95.57%

(2) Linear Discriminant Analysis

(a) Background

[0301] The commercial statistical package SPSS has procedures allowing simple linear discriminant functions to be design and tested.

[0302] A commonly used method is Fisher's Linear discriminant function. This finds the hyper-plane in feature space which gives a good separation of classes. For a two class problem where the class distributions have different means, but similar multivariate Gaussian distributions, this classifier gives optimum performance. The method can be extended heuristically to multi-class problems, but this was not applied in the study.

[0303] The method is simplistic in its approach but robust to problems associated with data sets containing a large number of features (the probes in our case number 27, giving problem for a data set comprising only some two hundred exemplars (cases)).

[0304] This package has a procedure for identifying the features which contribute well to the discrimination process. This "stepwise method" first finds the most discriminating feature. Other features are then sequentially added and evaluated against the classifier. Combinations are explored so the final solution may exclude features initially selected if better combinations are

found. The number of features is gradually increased until a statistical test shows the remaining features do not contribute reliably to the classification process.

[0305] An estimate of the performance is gained by using the leave one out method. This removes one sample from the data set to form the training set. The left out sample is retained as the test set, applied to the classifier, and the resulting classification accumulated in the confusion matrix. The procedure is repeated for case in the data. This procedure gives an unbiased estimate of performance, but the estimate will have a high variance.

[0306] In SPSS select the appropriate data set for analysis, select "Analyze", select "Classify", select "Discriminant...", on the table select "Fishers method", "leave one out testing" and "use stepwise method". Enter the diagnosis as the grouping variable and enter all the features as the independents. Enter 'OK" to complete the analysis. Pre-set values for other parameters were left as set.

[0307] The analysis output includes a list of the features used in the analysis, the canonical discriminant function and a confusion matrix and the correct-classification rate (1-error rate).

[0308] In order to compute an ROC curve the Canonical discriminant function is applied to the selected features to generate a new feature. In SPSS use Graphs, ROC to plot this curve.

ii. Hierarchical methods: Decision trees

(1) Background

[0309] Decision tree learning is one of the most widely used and practical methods for inductive inference. It is a method for classification that is robust to noisy data and capable of learning disjunctive expressions (Tom M. Mitchell, "Machine Learning", McGraw-Hill, New York, NY, 1997).

[0310] The most popular and accessible machine learning package is "C4.5" the source code of which is published in: (J. Ross Quinlan, "C4.5: Programs for Machine Learning", Morgan Kaufmann, San Mateo CA, 1993).

[0311] When a decision tree is being trained (on training data), the algorithm decides at each node of the tree which single attribute of the data to use at this node to best make a decision. Therefore when the tree is completely constructed, it will have selected some set of attributes to use and ignored others. In our application, using decision trees to process measurements gained from molecular probes, the decision tree has effectively chosen a panel of probes, and a method of combining the probe scores, which best explains the classification of the data. To obtain an

unbiased estimate of the panel performance, the resulting tree must be evaluated on data which was not used in the training. One standard technique for doing this is cross-validation. A 10-fold cross-validation was employed.

[0312] Cross-validation is a technique for making the very best use of limited data. In 10-fold cross-validation the data is randomly split into 10 nearly-equal sized partitions, taking care to have approximately the same number of cases in a class across each partition. Then, the decision tree is trained on partitions 2-9 combined and tested on partition 1, then trained on partitions 1,3-9 combined and tested on partition 2, and so on for 10 trials rotating the held-out test set through the data once. In this manner tests are only ever performed on held-out data and so are unbiased, and all data is tested exactly once so an aggregate error rate across the whole data set can be computed.

[0313] Trees are usually constructed until they are a very good fit to the training data, then they are "pruned" back by clipping off "noisy" branches and leaves. This improves the generalization ability of the decision tree on unseen data and is essential to obtain good performance. The C4.5 package includes two methods for pruning trees first a standard tree pruning algorithm, second a rule extraction algorithm. In general, the tree based method was found to give superior results on this data. Therefore, the rule-based method is not reported.

(2) Data Preparation

[0314] Data on the response of various probes to normal tissue and five different cancers (Adenocarcinoma, Large Cell Carcinoma, Mesothelioma, Small Cell Lung Cancer, and Squamous Cell Carcinoma) was obtained as described elsewhere. The H-scores for probes 1-28, and pathologists Pathologist 1 and Pathologist 2 were extracted from the database and put into a flat data file. For the decision tree analysis each data point (even by two pathologists on a same physical slide) was taken to be an independent observation of the effect of disease on staining. This may slightly positively bias the performance of classification but should have no effect on panel selection.

- The control categories of Emphysema, Granulomatous Disease, and Interstitial Lung Disease were grouped together and called "Normal".
- For the detection panel all the cancers were grouped together and called "Abnormal" making this a 2-class problem.

• For the single discrimination panel, the Normal cases were removed from the data to form a 5-class problem.

• For the hold-out discrimination panels, each cancer was held out in turn and the remaining cancers grouped into "Other" to give a set of five 2-class problems.

[0315] C4.5 requires a ".names" file which describes the data and the attributes to be included in the analysis. An example names file for the discrimination panel is, Table 26:

Table 26:

```
C4.5 Names file for MonoGen ZF21 diag data
Adenocarcinoma, Large Cell Carcinoma, Mesothelioma, Small Cell Lung Cancer,
Squamous Cell Carcinoma. | classes
        continuous.
        continuous.
       continuous.
        continuous.
        continuous.
        continuous.
        continuous.
       continuous.
        continuous.
        continuous.
P12 :
      continuous.
P13 :
       continuous.
        continuous.
P15 :
       continuous.
P16 :
        continuous.
P17. :
       continuous.
       ignore.
P18 :
P19 :
       continuous.
P20 :
       continuous.
P21 :
       continuous.
P22 :
        continuous.
P23 :
        continuous.
P24 :
        continuous.
P25 :
       continuous.
P26:
        continuous.
P27 :
        continuous.
P28 :
        continuous.
```

[0316] Probe 18 was missing from the data and was set to "ignore" in all the designs. Setting attributes to "ignore" in the names file is an easy and effective way of trimming probes from the panels and is used in the data analysis.

(3) Data Analysis

[0317] Ten-fold cross validation was run on each data set using the "kval.sh" script supplied with C4.5. Standard (default) parameters for the package were used. Cross validation is a technique developed for classifier training and testing on small data sets. It involves randomly splitting the data into N equal sized partitions. The classifier is then trained on N-1 partitions together and tested on the remaining partition. This is repeated N times.

[0318] Since the decision tree trained in one cross-validation(CV) trial may differ from the tree obtained in another (different in both probes selected, and tree coefficients) the number of times each probe was selected by the tree in 10 trials was computed.

[0319] The first cull of probes was done by setting to ignore any probe which did not occur in a pruned tree 5 or more times out of the 10 CV trials.

[0320] Then the cross-validation was repeated with this smaller set of candidate probes. The second cull of probes was done by setting to ignore any probe which did not occur in a pruned tree 5 or more times out of the 10 CV trials. If any further probes dropped out, a third CV run was done.

[0321] The panels were selected by the various runs, and their estimated error performance are shown in the results tables. The panel performance for decision tree analysis is shown below, in Table 27.

Table 27:

Panel Performance - Decision Trees

Detection Panel

Probes: 3, 7, 19, 25 and 28

Pair-wise Discrimination 4, 6, 14, 19 and 23

Pair-wise Discrimination 3, 6, 17, 19 and 25

Pair-wise Discrimination 1, 5, 10, 13, 21, 27 and 28

Pair-wise Discrimination 3, 12 and 16

Pair-wise Discrimination 12, 17, 20, 23 and 25

Detection (without probe 7) 6, 10, 16 and 19

Detection (only commercially preferred probes) 5, 6, 10, 16, 19 and 23

同學所屬第28 8	Cancer	Control
Cancer	99.42%	0.58%
Control .	17.82%	82.18%

为是分别数 。	Adeno	Others
Adeno	67.74%	32.26%
Others	11.20%	88.80%

建筑等于企业的	Squamous	Others
Squamous	70.59%	29.41%
Others .	4.07%	95.93%

a. Included the	Large Cell	Others
Large Cell :	36.36%	63.64%
Others	7.37%	92.63%

2005年第993	Mesothelioma	Others
Mesothelioma	82.05%	17.95%
Others	5.00%	95.00%

	Small Cell	Others ·
Small Cell	69.23%	30.77%
Others .	1.49%	98.51%

多数数值的 多数	Cancer	Control
Cancer	89.60%	10.40%
Control	3.30%	96.70%

ENTER SERVICE	Cancer	Control
Cancer	92.80%	7.20%
Control	5.49%	94.51%

[0322] An example decision tree structure is shown in below, in Tables 28 and 29, for discriminating between Small Cell Lung Cancer and the remaining four types of cancer.

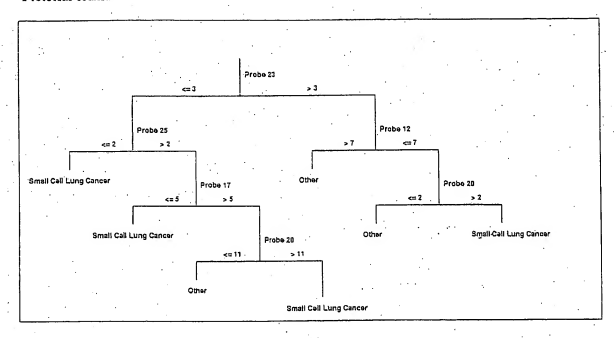
Table 28:

C4.5 output format:

```
P23 <= 3:
 P25 <= 2 : Small Cell Lung Cancer (18.0)
 P25 > 2:
 | P17 <= 5 : Small Cell Lung Cancer (2.0)
  | P17 > 5:
  | P20 \le 11 : Other (9.0)
P20 > 11 : Small Cell Lung Cancer (2.0)
P23 > 3:
P12 > 7: Other (120.0)
P12 <= 7:
| P20 \le 2 : Other (5.0)
P20 > 2 : Small Cell Lung Cancer (4.0)
Tree saved
Evaluation on training data (160 items):
     Before Pruning
                         After Pruning
                          Errors Estimate
    Size
           Errors Size
     13
          0(0.0%)
                    13
                         0(0.0%) (5.2%)
```

Table 29:

Pictorial format:



[0323] The panel performance for stepwise linear discriminant is shown below, in Table 30:

Table 30:

Panel Performance - Stepwise LD

Detection Panel 1, 4, 7, 15 and 16

Pair-wise Discrimination 4, 5, 14, 19, 20, 25 and 27

Pair-wise Discrimination 1, 2, 3, 24, 25 and 26

Pair-wise Discrimination 1 and 7

Pair-wise Discrimination 3, 12 and 16

Pair-wise Discrimination 12, 19, 22 and 23

Detection (without probe 7)
1, 2, 3, 4, 10, 11, 15, 16, 23, 24, 27 and 28

Detection (only commercially preferred probes) 8, 10, 11, 19, 23 and 28

PER NAME OF THE	Cancer		Control
Cancer .		92.24%	7.76%
Control		1.16%	98.84%

Line II all the	Adeno		Others
Adeno	· ·	91.67%	8.33%
Others		5.43%	94.57%

2000年1月1日	Squamous	Others
Squamous	88.00%	12.00%
Others	6.59%	93.41%

可以类型 (200)	Large Cell	Others
Large Cell	80.95%	19.05%
Others	26.32%	73.68%

	Mesothelioma	Others
Mesothelioma	96.67%	3.33%
Others ·	4.65%	95.35%

MASSACTION OF	Small Cell	Others
Small Cell	93.75%	6.25%
Others	. 5.00%	95.00%

PROPERTY OF STATE	Cancer	Control
Cancer	85.34%	14.66%
Control	2.33%	97.67%

REST WORKS	Cancer		Control
Cancer		81.20%	18.80%
Control		1.16%	98.84%

[0324] The panel performance for stepwise logistic regression analysis is shown below, in Table 31:

Table 31:

Panel Performance - Stepwise LR

Detection Panel 6, 7, 12, 23 and 25

Pair-wise Discrimination 14, 19, 20, 25 and 27

Pair-wise Discrimination 3 and 10

Pair-wise Discrimination 1, 4, 6, 16 and 21

Pair-wise Discrimination 3, 7, 12 and 16

Pair-wise Discrimination 12, 13 and 23

Detection (without probe 7) 1, 10, 19, 23 and 28

Detection (only commercially preferred probes) 10, 19, 20, 23 and 28

	Cancer	Control
Cancer	97.49%	2.63%
Control	2.51%	97.49%

是经历的特殊	Adeno	Others
Adeno	96.39%	3.61%
Others	12.29%	87.71%

国际发生的对象	Squamous	Others
Squamous	94.93%	5.07%
Others	35.86%	64.14%

THE PERSON NAMED IN	Large Cell	Others
Large Cell	95.11%	4.89%
Others	61.00%	39.00%

经工业的现在	Mesothelioma	Others
Mesothelioma	95.07%	4.93%
Others	10.89%	89.11%

	Small Cell	Others
Small Cell	98.90%	1.10%
Others	4.00%	96.00%

深度 计多数数字	Cancer	Control
Cancer	94.00%	6.00%
Control	5.80%	94.20%

	Cancer	Control
Cancer	93.88%	6.12%
Control	6.39%	93.61%

iii. Neural networks and alternative methods

[0325] Artificial neural networks ANN's are candidate pattern recognition techniques which could readily be applied to select features and design classifiers in association with this invention. However such techniques give little insight to the structure of the data and the influence of particular probes in the way that LDF gives. For this reason this class of algorithm was not used in this study. LDF stands for linear discriminant function, a linear combination of features whose result is thresholded to determine the classification.

[0326] This class of techniques includes algorithms such as Multi-Layer Perceptron MLP, Back-Prop, Kohonen's Self-Organizing Maps, Learning Vector Quantization, K-nearest neighbors and Genetic Algorithms.

iv. Special topics

(1) Assumptions

- Linear discriminant analysis
 - Assumes the covariance matrices for the two classes are equal.
 - Minimizes the cost of misclassification only when the two classes are multivariate normal.
 - Assumes that the explanatory variables are continuous rather than categorical (in this study, the H-scores are categorical while in practice (i.e., in an automated system) intensity can be measured on a continuous scale).
- Logistic regression (binomial generalized linear models)

[0327] See Venerables and Ripley, chapter 7 ("Modern Applied Statistics with S-PLUS" (W.N. Venables and B.D. Ripley, Springer-Verlag, New York, 1999)).

(2) Marker rejection (de-selection)

[0328] Computerized implementations of discriminant analysis and regression procedures include stepwise variable selection procedures; e.g., stepAIC in R. These procedures are designed to select the *best* subset of variables for use as explanatory variables. In reality, because of the step-by-step nature of these procedures, there is no guarantee that the best variables are selected for prediction (Johnson and Wichern, p. 299). Nevertheless such procedures do provide the basis for marker selection and de-selection.

(3) Pairwise tests

[0329] Inherent problems in designing multiclass classifiers is discussed in "Applied Mulitvariate Statistical Analysis", R. A. Johnson and D. W. Wichern, 2nd Ed,1988, Prentice-Hall, N.J. This is motivation for developing several separate two-class classifiers (discrimination panel).

(4) Redundancy consideration in panel composition

[0330] "Linear models form the core of classical statistics and are still the basis of much of statistical practice" "Modern Applied Statistics with S-PLUS" (W.N. Venables and B.D. Ripley, Springer-Verlag, New York, 1999). Linear models are the foundation for the t-test,

analysis of variance (ANOVA), regression analysis, as well as a variety of multivariate methods including discriminant analysis. Explanatory variables may or may not enter the model as first-order terms. This is true also of (non-linear) logistic regression. The logistic regression model is simply a non-linear transformation of the linear regression model: the dependent variable is replaced by a log odds ratio (logit). In summary these statistical methods are based on linear relationships between the explanatory variables. Consequently, one avenue for seeking redundancy in panels is to identify highly correlated variables (markers). It may be possible to replace one marker with the other in a panel to achieve similar performance.

[0331] Another avenue for seeking redundancy in panels is to undertake a "best subsets" regression analysis. Given a starting model with all of the explanatory variables of interest, the aim is to find the best single-variable regression models, the best two-variable regression, etc. This methodology is implemented in the SAS statistical package.

(5) Use of weighting scores

(a) Commercial and clinical considerations

[0332] For many reasons, including strategic and commercial factors; cost; availability; ease of use, it may be preferred to encourage the selection of certain probes in a panel and penalize the selection of others, at the same time trading this off against panel size or performance.

(b) Attribute costing

[0333] Methods for such attribute weighting (in decision trees) have been proposed in the machine learning literature in other contexts such as the incorporation of background knowledge (M. Nunez, "The Use of Background Knowledge", Machine Learning 6: 231-250, 1991.), and the differential cost of obtaining information from robotic sensors (M. Tan, "Cost-sensitive Learning of Classification Knowledge and its Applications in Robotics", Machine Learning. 13: 7-33, 1993).

[0334] Both of these cost-sensitive algorithms have been implemented in the literature by minor changes to the standard machine learning software package known as "C4.5 (J. Ross Quinlan, "C4.5: programs for machine learning", Morgan Kaufmann, CA. 1993). For convenience, this approach was followed to implement the "EG2" algorithm of Nunez.

[0335] In the C4.5 decision tree construction phase, the algorithm compares each available attribute to split on and chooses the single one which maximizes the information gain, Gi. In the

EG2 algorithm, $(2^{Gi}-1)/(Ci+1)$ is maximized which incorporates the cost of information for attribute i, Ci. The vector of weights need to be set a prior by the user.

(i) Code Modifications

[0336] The C4.5 source code was modified to implement the economic generalizer "EG2" algorithm proposed by M. Nunez (The Use of Background Knowledge, Machine Learning 6: 231-250, 1991).

[0337] The exact modifications to the C4.5 package are as follows. After the following lines in file "R8/Src/contin.c". (J. Ross Quinlan, "C4.5: programs for machine learning", Morgan Kaufmann, CA. 1993).

```
ForEach(i, Xp, Lp - 1)
{
    if ( (Val = SplitGain[i] - ThreshCost) > BestVal )
    {
        BestI = i;
        BestVal = Val;
    }
}
```

The new line:

```
BestVal = (powf(2.0, BestVal) - 1.0) / (AttributeCosts(Att) + 1.0);
```

is inserted. Where the vector of attribute costs has been previously read in from a text file maintained by the user.

(ii) Experimental Methodology.

[0338] The commercially preferred probes are: 2,4,5,6,8,10,11,12,16,19,20,22,23,28.

[0339] For the sake of example, suppose the above probes are commercially preferred due to cost and it is desired to reselect the detection panel taking this cost into account.

[0340] The modified C4.5 decision tree software was used to give the commercially preferred probes a penalty of zero and non-commercially preferred probes a penalty of two. The 10-fold cross validated panel selection methodology (as described elsewhere) was run using the modified C4.5 algorithm.

(iii) Results

[0341] The standard decision tree detection panel consists of probes 3, 7, 19, 25, 28. Resulting Panel Members: are 2, 6, 7, 10, 19, 25, 28 which used only 2 commercially preferred probes, P7 and P25. Note these probes have been selected by the method in spite of their increased cost due

to their superior performance on this data. The panel is now larger: 7 probes versus 5 originally. There is no demonstratable drop in panel performance on this data although the performance will now be sub-optimal as a trade off against the reduced cost of probes.

(iv) Conclusion

[0342] A straightforward way has been established for incorporating costs of using probes into the panel selection methodology.

(c) Misclassification costing

(i) Background

[0343] For many reasons it may be desired to select an optimal panel bearing in mind that the costs of the different kinds of classification errors may vary. For example, it may be desired to select a panel which has an increased sensitivity to one disease (say Large Cell Carcinoma) and be willing to trade this off against reduced specificity and sensitivity elsewhere in the confusion matrix.

[0344] In theory a matrix of misclassification costs (of the same dimensions as the confusion matrix) to incorporate all the possible combinations of costs may be needed. In practice, only those costs which are non unity (the default) are entered.

[0345] The commercial decision tree software See5. (RuleQuest Research Pty Ltd, 30 Athena Avenue, St Ives Pathologist 3SW 2075, Australia. (http://www.rulequest.com)) incorporates this capability and was used in the following demonstration.

(ii) Aim

[0346] The standard joint discrimination panel (described elsewhere) consists of the members: P2, 3, 4, 5, 12, 14, 16, 19, 22, 23, 28. And gives the following estimated confusion matrix:

	(a).	(b)	(c)	(d)	(e)	<-classified as	
	24	4	2	5	2	(a): class Adenocarcinoma	
•	. 8	7	3	5	. 4	(b): class Large Cell Carcinoma	
	1	1	33	1	4	(c): class Mesothelioma	
	.6	2	1	. 23		(d): class Small Cell Lung Cancer	
	4	4	3	2	· 24		• .

[0347] The sensitivity of Large Cell Carcinoma is low at 26 percent. If one wished to increase this sensitivity in a newly designed panel, the following method may be employed.

(iii) Methodology

[0348] The following costs file was generated:

```
| costs file for ZF21Discrim
| Increase sensitivity for "Large Cell Carcinoma"
| Mesothelioma, Large Cell Carcinoma: 10
Adenocarcinoma, Large Cell Carcinoma: 10
Mesothelioma, Large Cell Carcinoma: 10
Small Cell Lung Cancer, Large Cell Carcinoma: 10
Squamous Cell Carcinoma, Large Cell Carcinoma: 10
```

[0349] This file upweights the misclassification of Large Cell Carcinoma as any of the other cancers by a factor of 10. This will tend to increase the sensitivity of detection in this class (with reduced performance elsewhere) but no weighting can ensure perfect classification.

[0350] The standard decision tree panel selection methodology was applied (using See5 instead of C4.5).

(iv) Results

[0351] The new panel members are: P2, 3, 4, 5, 6, 9, 12, 14, 16, 17, 25, 28. With an estimated performance of:

	(a)	(a)	(c)	(d)	(e)	<-classified as
	20 3 1 · 2	13 13 9 9	1 3 27	1 2 2 21 1	2 6 1	 (a): class Adenocarcinoma (b): class Large Cell Carcinoma (c): class Mesothelioma (d): class Small Cell Lung Cancer (e): class Squamous Cell
Carcinoma			•			

[0352] The above demonstrates that the estimated sensitivity of Large Cell Carcinoma has now increased to 48%.

(v) Conclusion

[0353] A straightforward way has been demonstrated for incorporating the differential costs of misclassification into the panel selection methodology.

d. Performance metrics

[0354] Outputs provided by the analysis indicating the estimated performance of each method include:

i. ROC analyses

[0355] Receiver Operating Characteristic (ROC) curves show the estimated percentage (or per unit probability) of false positive and false negative scores for different threshold levels in the classifier. An indifferent classifier, unable to discriminate better than random choice, would present a ROC curve with equal true and false readings. The area under this curve would be 50% (0.5 probability).

[0356] Area Under the Curve (AUC) is often used as an overall estimate of classifier performance and most commercial discriminant function packages compute this figure. A perfect classifier would have 100% Area Under the Curve, a useless classifier would have an AUC near 50% (0.5 probability).

ii. Confusion matrices: counts and percentages

[0357] Confusion matrices show how data from the test set was classified. For pair wise tests these are counts of true positive, false positive, true negative or false negative scores. These may be shown as actual counts or as percentages. For the multi-way Panel, which attempts to give a unique diagnosis with one panel only, the confusion matrix would show counts for each correct classification. For instance, each time Small Cell carcinoma is detected as such it would be entered in one diagonal of the matrix. Incorrect scores; for instance, how often a small cell carcinoma is incorrectly identified as squamous cell cancer would be entered in the appropriate off-diagonal element of the matrix. Error Rates are used to summarize data in the confusion matrix as the sum of all false classifications divided by the total number of classifications made, expressed as a percentage.

iii. Sensitivity and specificity

[0358] Specificity refers to the extent to which any definition excludes invalid cases. If a definition has poor specificity, it is high in false positives. This means that it labels individuals as having a disorder when there is really no disorder present. Sensitivity refers to the extent to which any definition includes all valid cases. If a definition has poor sensitivity, it is high in false negatives (individuals who have a disorder present are falsely being diagnosed as not having one).

3. Data analysis and results

- a. Sample size and variability
- Of the 354 cases in the combined Pathologist 1 and Pathologist 2 data set, only 202 cases possessed an H score for every marker (variable or feature).
- The small number of complete observations and the large number of variables leads to estimation problems (curse of dimensionality). Hence it is necessary to prune severely back the number of variables used to build a classifier.
- Due to the small number of observations it is not prudent to divide the data into separate training and testing sets (necessary for the robust estimation of classifier performance). For this reason, it was necessary to use resampling methods (such as cross-validation and multiple random trials).
- The design of a multiclass classifier for cancer discrimination is difficult because there are so few observations for each type of cancer.

b. De-selected markers

[0359] Markers were de-selected using the methodology described above. Markers that were de-selected are represented by non-selection in the panels.

- c. Detection panel(s) composition
 - i. Selected marker probes

[0360] The selected marker probes for all three methods are summarized in Figure 5.

ii. Minimum selected marker set

[0361] For the detection panel it is clear that probe 7 delivered the best detection performance for a single marker. Combinations of probes were analyzed to see if a reliable panel could be obtained with more probes.

(1) Method

[0362] The Logistic Regression method allows best subsets to be ranked in terms of a performance measure (Fisher'score). This analysis was used to select the combinations from 1 through 5 probes. Fishers linear discriminant function and logit models (logistic regression) were used to illustrate the performance of these combinations. Data shown above.

(2) Conclusions

[0363] Probe 7 performs well on its own as a classifier; however, a drawback to using probe 7 alone is that probe 7 has a high false negative score. The best performance using Fishers linear

discriminant function as a classifier was with probes 7 and 16. The variability of results amongst panels using other combinations suggests the noise added by more features is outweighing any potential to improve classification scores. The small number of incorrectly scored samples gives a poor representation of the statistics of these rarer events. A classifier designed with a larger number of cases may allow a better classifier to be designed. Techniques to select best combinations of probes using different classifiers may produce a different best panel, depending on the structure of the data.

iii. Supplemental markers

[0364] It is shown that panels can be designed to suit the availability of different probes. Different methodologies can be used for selecting these subsets: Decision Trees, Logistic Regression, and Linear Discriminant Functions. Data are shown above.

[0365] Using SPSS a Fisher's Linear Discriminant function was applied to the scores obtained from the panel in which constrains were applied due to access constraints. For example, all of the probes come from one vendor. Again, the stepwise option was selected to find the best combination of features. Performance was estimated using the Leave-One-Out cross validation test.

iv. Alternative markers: biological mechanisms of action (functionally equivalent markers)

[0366] A person of ordinary skill in the art is able to determine functionally equivalent markers. The functional behaviors of the markers used in the panel are described throughout this document.

v. Marker localization

[0367] The localizations of the various markers used in this study are described elsewhere in this document.

vi. Panel Performance

[0368] The performance of the three methods is shown above.

- vii. Limitations on interpretation of panel performance
- Due to small data set and the need to employ resampling methods, there is the danger that the classifiers have been over-trained (made to fit the data too closely).
- The panel performance using cytology specimens is difficult to forecast accurately since it is not clear whether sputum cytology samples will contain adequate numbers of cells

that are representative of the cells analyzed in the histological validation studies.

Nevertheless, given an adequate cellular sample size, one would expect the optimized panel to behave similarly with cytological specimens.

- d. Discriminant Panel Composition
 - i. A single 5-way panel for all cancers

[0369] Of the three analysis techniques, only a decision tree is amenable to a single 5-way panel. A single decision tree was therefore constructed to simultaneously classify all types of lung cancer. The panel members are shown Figure 5. The panel performance is shown above in the panel performance tables.

ii. Panels for discriminating a single type of lung cancer against all others

[0370] Linear discriminant functions are not well suited to performing simultaneous multi-class discrimination. The performance of five separate classifiers, each designed separately to discriminate one of the cancers from a pooled set of all the cancers, was analyzed. Such combinations have the potential to classify none of the cases as having one of the candidate cancers, or classify a single case as having two or more of the candidate cancers. This has a potential advantage in identifying inconsistent cases for further review.

[0371] It has been seen that the overall error rate of a single discriminant panel for all cancer types has a fairly high error rate (a five way classifier). In the panel performance data shown above, the performance of five pair-wise classifiers, each designed to identify one cancer from the four other possible cancers is shown. This approach is amenable to analysis using Decision Trees, and Linear Discriminant functions. The technique has the potential to deliver an ambiguous finding when applied, giving two or more diagnoses for a single patient, suggesting further clinical investigation. The technique has the potential to deliver no finding, again suggesting further investigation (perhaps a re-test with the detection panel).

iii. Panels to account for possibility of false positive cases from detection panels

[0372] A further panel can be trained to discriminate among the false positive cases (from the detection panel) and the five cancer types. This involves selecting those individual cases from the detection panel that were incorrectly classified as abnormal. This trains a dedicated classifier on the 'harder' problem of detecting these 'special' cases. However, while this is a theoretically

sound task, the data set only yielded four of these cases and the population was deemed to be under-represented for analysis.

iv. Selected Markers

[0373] The selected marker probes for all three methods are summarized in Figure 5.

v. Minimum selected marker set

[0374] This topic is addressed below under "Robustness of Approach Demonstrated by Similar Results Using Different Methods."

vi. Supplemental markers

[0375] This topic is addressed below under "Robustness of Approach Demonstrated by Similar Results Using Different Methods."

vii. Alternative markers: biological mechanisms of action
[0376] A person of ordinary skill in the art is able to determine functionally equivalent markers. The functional behaviors of the markers used in the panel are described throughout this document.

viii. Marker localization

[0377] The localization of the various markers used in this study are described throughout this document.

ix. Panel Performance

[0378] The performance of the three methods is summarized in Figure 5.

e. Effect of weighting parameters

[0379] In addition to user-supplied weighting criteria for markers and also for disease states (classes) as discussed earlier, one can also use a binary weighting scheme. For example, if all non-DAKO supplied probes are weighted "0" and all DAKO-supplied probes are weighted "1", then the optimized panel will contain only DAKO-supplied probes. This is an important product design capability for any vendor who intends to develop and market molecular diagnostic panel kits using only their supplies.

f. Effect of using other (non H-score) objective scoring parametersi. Background

[0380] The Pathology Review sheet contains a set of boxes as follows, in Table 32: Table 32:

Intensity	None	Weak	Moderate	Intense
0-5%	□0	□ 0	□ 0	□0 .
6-25%	01	□ 1		
26-50%	□ 2	□ 2	□ 2	□ 2
51-75%	-□ 3	□ 3	□ 3	3
>75%	□4	□4	·D 4	□ 4

[0381] The standard scoring system uses the "H score" which is obtained by grading the intensity as: none = 0, weak = 1, moderate = 2, intense = 3, and the percentage cells as: 0-5% = 0, 6-25% = 1, 26-50% = 2, 51-75% = 3, >75% = 4, and then multiplying the two grades together. For example, 50% weakly stained plus 50% moderate stained would score $10 = 2 \times 2 + 2 \times 3$.

ii. Method

[0382] An alternative scoring method was analyzed in which the response was divided into low, medium and high as follows:

- (a) if more than 50% of cells had moderate or above stain → HIGH
- (b) if more than 50% of cells had no stain → LOW
- (c) otherwise → MEDIUM

[0383] The decision tree detection panel selection methodology was repeated using this 3-level factor instead of H-score. This caused the tree to split into 3 branches at each node, if required.

iii. Results

[0384] The panel selected was: Probes 3, 7, 10, 11, 16, 19, 20, 28. With an estimated performance of:

Classified as ->		(a)	(b)	
<u> </u>	Control (a)	79	22	Specificity = 78%
·.	Cancer (b)	24	149	Sensitivity = 86%

This should be compared to the reference performance with H-scores of:

Classified as ->	(a)	.(p)	
Control (a)	85	6.	Specificity = 93%
Cancer (b)	5	120	Sensitivity = 96%

iv. Conclusions

- There is a substantial loss of performance (larger panels, lower sensitivity and lower specificity) when the proposed alternative scoring system is used.
- Treating the H-score as a continuous variable (in the range 0 to 12) seems to be near optimal for panel selection on the data examined.
- The many other possible scoring systems have not been examined, but may be feasible
 and applicable to the experimentally tested panel design and development methodology.

4. Lung Cancer Detection and Discrimination Panels

[0385] Listed below are exemplary lung cancer detection and discrimination panels determined by the above illustrative example. It is noted that although the panels listed below recite specific probes, each specific probe may be substituted by a correlate probe or a functionally related probe.

Detection (No Constraints)

- anti-Cyclin A combined with one or more additional probes.
- anti-Cyclin A, anti-human epithelial related antigen (MOC-31).
- anti-Cyclin A, anti-ER-related P29.
- anti-Cyclin A, anti-mature surfactant apoprotein B.
- anti-Cyclin A, anti-human epithelial related antigen (MOC-31), anti-VEGF.

• anti-Cyclin A, anti-human epithelial related antigen (MOC-31), anti-mature surfactant apoprotein B.

- anti-Cyclin A, anti-mature surfactant apoprotein B, anti-human epithelial related antigen (MOC-31), anti-VEGF.
- anti-Cyclin A, anti-mature surfactant apoprotein B, anti-human epithelial related antigen (MOC-31), anti-surfactant apoprotein A.
- anti-Cyclin A, anti-mature surfactant apoprotein B, anti-human epithelial related antigen (MOC-31), anti-VEGF, anti-surfactant apoprotein A.
- anti-Cyclin A, anti-mature surfactant apoprotein B, anti-human epithelial related antigen (MOC-31), anti-VEGF, anti-Cyclin D1.
- anti-Cyclin A, anti-human epithelial related antigen (MOC-31) combined with one or more additional probes.
- anti-Cyclin A, anti-ER-related P29 combined with one or more additional probes.
- anti-Cyclin A, anti-mature surfactant apoprotein B combined with one or more additional probes.
- anti-Cyclin A, anti-human epithelial related antigen (MOC-31), anti-VEGF combined with one or more additional probes.
- anti-Cyclin A, anti-human epithelial related antigen (MOC-31), anti-mature surfactant apoprotein B combined with one or more additional probes.
- anti-Cyclin A, anti-mature surfactant apoprotein B, anti-human epithelial related antigen (MOC-31), anti-VEGF combined with one or more additional probes.
- anti-Cyclin A, anti-mature surfactant apoprotein B, anti-human epithelial related antigen (MOC-31), anti-surfactant apoprotein A combined with one or more additional probes.
- anti-Cyclin A, anti-mature surfactant apoprotein B, anti-human epithelial related antigen (MOC-31), anti-VEGF, anti-surfactant apoprotein A combined with one or more additional probes.
- anti-Cyclin A, anti-mature surfactant apoprotein B, anti-human epithelial related antigen (MOC-31), anti-VEGF, anti-Cyclin D1 combined with one or more additional probes.

Detection (W/O anti-Cyclin A)

• anti-Ki-67 combined with one or more additional probes.

- anti-Ki-67 combined with any one probe selected from the group consisting of anti-VEGF, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen and anti-mature surfactant apoprotein B.
- anti-Ki-67 combined with any two probes selected from the group consisting of anti-VEGF, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen and anti-mature surfactant apoprotein B.
- anti-Ki-67 combined with any three probes selected from the group consisting of anti-VEGF, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen and anti-mature surfactant apoprotein B.
- anti-Ki-67 combined with any four probes selected from the group consisting of anti-VEGF, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen and anti-mature surfactant apoprotein B.
- anti-Ki-67 combined with any five probes selected from the group consisting of anti-VEGF, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen and anti-mature surfactant apoprotein B.
- anti-Ki-67, anti-VEGF, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen and anti-mature surfactant apoprotein B.
- anti-Ki-67 combined with any one probe selected from the group consisting of anti-VEGF, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen and anti-mature surfactant apoprotein B, and with one or more additional probes.
- anti-Ki-67 combined with any two probes selected from the group consisting of anti-VEGF, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen and anti-mature surfactant apoprotein B, and with one or more additional probes.
- anti-Ki-67 combined with any three probes selected from the group consisting of anti-VEGF, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen and anti-mature surfactant apoprotein B, and with one or more additional probes.

• anti-Ki-67 combined with any four probes selected from the group consisting of anti-VEGF, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen and anti-mature surfactant apoprotein B, and with one or more additional probes.

- anti-Ki-67 combined with any five probes selected from the group consisting of anti-VEGF, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen and anti-mature surfactant apoprotein B, and with one or more additional probes.
- anti-Ki-67, anti-VEGF, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen, anti-mature surfactant apoprotein B and one or more additional probes.

Detection With Commerically Preferred Probes

- anti-Ki-67 combined with one or more additional probes.
- anti-TTF-1 combined with one or more additional probes.
- anti-EGFR combined with one or more additional probes.
- anti-proliferating cell nuclear antigen combined with one or more additional probes.
- two probes selected from the group consisting of anti-Ki-67, anti-TTF-1, anti-EGFR and anti-proliferating cell nuclear antigen.
- three probes selected from the group consisting of anti-Ki-67, anti-TTF-1, anti-EGFR and anti-proliferating cell nuclear antigen.
- anti-Ki-67, anti-TTF-1, anti-EGFR and anti-proliferating cell nuclear antigen.
- two probes selected from the group consisting of anti-Ki-67, anti-TTF-1, anti-EGFR and anti-proliferating cell nuclear antigen, and one or more additional probes.
- three probes selected from the group consisting of anti-Ki-67, anti-TTF-1, anti-EGFR and anti-proliferating cell nuclear antigen, and one or more additional probes.
- anti-Ki-67, anti-TTF-1, anti-EGFR, anti-proliferating cell nuclear antigen, and one or more additional probes.

Discrimination Between Adenocarcinoma And Other Lung Cancers

• anti-mucin 1 and anti-TTF-1.

anti-mucin 1 and anti-TTF-1 combined with any one probe selected from the group consisting of anti-VEGF, anti-surfactant apoprotein A, anti-BCL2, anti-ER-related P29 and anti-Glut 3.

- anti-mucin 1 and anti-TTF-1 combined with and two probes selected from the group consisting of anti-VEGF, anti-surfactant apoprotein A, anti-BCL2, anti-ER-related P29 and anti-Glut 3.
- anti-mucin 1 and anti-TTF-1 combined with any three probes selected from the group consisting of anti-VEGF, anti-surfactant apoprotein A, anti-BCL2, anti-ER-related P29 and anti-Glut 3.
- anti-mucin 1 and anti-TTF-1 combined with any four probes selected from the group consisting of anti-VEGF, anti-surfactant apoprotein A, anti-BCL2, anti-ER-related P29 and anti-Glut 3.
- anti-VEGF, anti-surfactant apoprotein A, anti-mucin 1, anti-TTF-1, anti-BCL2, anti-ER-related P29 and anti-Glut 3.
- anti-mucin 1, anti-TTF-1 and one or more additional probes.
- anti-mucin 1 and anti-TTF-1 combined with any one probe selected from the group consisting of anti-VEGF, anti-surfactant apoprotein A, anti-BCL2, anti-ER-related P29 and anti-Glut 3, and with one or more additional probes.
- anti-mucin 1 and anti-TTF-1 combined with and two probes selected from the group consisting of anti-VEGF, anti-surfactant apoprotein A, anti-BCL2, anti-ER-related P29 and anti-Glut 3, and with one or more additional probes.
- anti-mucin 1 and anti-TTF-1 combined with any three probes selected from the group consisting of anti-VEGF, anti-surfactant apoprotein A, anti-BCL2, anti-ER-related P29 and anti-Glut 3, and with one or more additional probes.
- anti-mucin 1 and anti-TTF-1 combined with any four probes selected from the group consisting of anti-VEGF, anti-surfactant apoprotein A, anti-BCL2, anti-ER-related P29 and anti-Glut 3, and with one or more additional probes.
- anti-VEGF, anti-surfactant apoprotein A, anti-mucin 1, anti-TTF-1, anti-BCL2, anti-ER-related P29, anti-Glut 3 and one or more additional probes.

Discrimination Between Squamous Cell Carcinoma And Other Lung Cancers

- anti-CD44v6 combined with one or more additional probes.
- anti-CD44v6 combined with any one probe selected from the group consisting of anti-VEGF, anti-thrombomodulin, anti-Glut 1, anti-ER-related P29 and anti-melanoma-associated antigen 3.
- anti-CD44v6 combined with any two probes selected from the group consisting of anti-VEGF, anti-thrombomodulin, anti-Glut 1, anti-ER-related P29 and anti-melanomaassociated antigen 3.
- anti-CD44v6 combined with any three probes selected from the group consisting of anti-VEGF, anti-thrombomodulin, anti-Glut 1, anti-ER-related P29 and anti-melanoma-associated antigen 3.
- anti-CD44v6 combined with any four probes selected from the group consisting of anti-VEGF, anti-thrombomodulin, anti-Glut 1, anti-ER-related P29 and anti-melanoma-associated antigen 3.
- anti-CD44v6, anti-VEGF, anti-thrombomodulin, anti-Glut 1, anti-ER-related P29 and anti-melanoma-associated antigen 3.
- anti-CD44v6 combined with any one probe selected from the group consisting of anti-VEGF, anti-thrombomodulin, anti-Glut 1, anti-ER-related P29 and anti-melanoma-associated antigen 3, and with one or more additional probes.
- anti-CD44v6 combined with any two probes selected from the group consisting of anti-VEGF, anti-thrombomodulin, anti-Glut 1, anti-ER-related P29 and anti-melanoma-associated antigen 3, and with one or more additional probes.
- anti-CD44v6 combined with any three probes selected from the group consisting of anti-VEGF, anti-thrombomodulin, anti-Glut 1, anti-ER-related P29 and anti-melanoma-associated antigen 3, and with one or more additional probes.
- anti-CD44v6 combined with any four probes selected from the group consisting of anti-VEGF, anti-thrombomodulin, anti-Glut 1, anti-ER-related P29 and anti-melanoma-associated antigen 3, and with one or more additional probes.
- anti-CD44v6, anti-VEGF, anti-thrombomodulin, anti-Glut 1, anti-ER-related P29, anti-melanoma-associated antigen 3 and one or more additional probes.

Discrimination Between Large Cell Carcinoma And Other Lung Cancers

- anti-VEGF combined with one or more additional probes.
- anti-VEGF and anti-p120.
- anti-VEGF and anti-Glut 3.
- anti-VEGF, anti-p120 and anti-Cyclin A.
- anti-VEGF, anti-p120 and one or more additional probes.
- anti-VEGF, anti-Glut 3 and one or more additional probes.
- anti-VEGF, anti-p120, anti-Cyclin A and one or more additional probes.

Discrimination Between Mesothelioma And Other Lung Cancers

- anti-CD44v6 combined with one or more additional probes.
- anti-proliferating cell nuclear antigen combined with one or more additional probes.
- anti-human epithelial related antigen (MOC-31) combined with one or more additional probes.
- two probes selected from the group consisting of anti-CD44v6, anti-proliferating cell nuclear antigen and anti-human epithelial related antigen (MOC-31), combined with one or more additional probes.
- anti-CD44v6, anti-proliferating cell nuclear antigen, anti-human epithelial related antigen (MOC-31) and one or more additional probes.

Discrimination Between Small Cell And Other Lung Cancers

- anti-proliferating cell nuclear antigen combined with one or more additional probes.
- anti-BCL2 combined with one or more additional probes.
- anti-EGFR combined with one or more additional probes.
- two probes selected from the group consisting of anti-proliferating cell nuclear antigen, anti-BCL2 and anti-EGFR.
- anti-proliferating cell nuclear antigen, anti-BCL2, anti-EGFR.
- two probes selected from the group consisting of anti-proliferating cell nuclear antigen, anti-BCL2 and anti-EGFR, combined with one or more additional probes.
- anti-proliferating cell nuclear antigen, anti-BCL2, anti-EGFR and one or more additional probes.

Simultaneous Discrimination Of Adenocarcinoma, Squamous Cell Carcinoma, Large Cell Carcinoma, Mesothelioma And Small Cell Carcinoma

- two or more probes selected from anti-VEGF, anti-thrombomodulin, anti-CD44v6, anti-surfactant apoprotein A, anti-proliferating cell nuclear antigen, anti-mucin 1, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-N-cadherin, anti-EGFR and anti-proliferating cell nuclear antigen.
- anti-VEGF, anti-thrombomodulin, anti-CD44v6, anti-surfactant apoprotein A, anti-proliferating cell nuclear antigen, anti-mucin 1, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-N-cadherin, anti-EGFR and anti-proliferating cell nuclear antigen.
- two or more probes selected from anti-VEGF, anti-thrombomodulin, anti-CD44v6, anti-surfactant apoprotein A, anti-proliferating cell nuclear antigen, anti-mucin 1, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-N-cadherin, anti-EGFR and anti-proliferating cell nuclear antigen, combined with one or more additional probes.
- anti-VEGF, anti-thrombomodulin, anti-CD44v6, anti-surfactant apoprotein A, anti-proliferating cell nuclear antigen, anti-mucin 1, anti-human epithelial related antigen (MOC-31), anti-TTF-1, anti-N-cadherin, anti-EGFR and anti-proliferating cell nuclear antigen, combined with one or more additional probes.

5. Conclusions

- a. Validity of panel approach to molecular diagnostics
 - i. Non-intuitive solutions

[0386] Histograms were plotted (PathologistData.xls, worksheet: Histograms) showing the distribution of marker scores for each probe for Control vs. Cancer. It is clear from these histograms that an intuitive selection of probes for specific panels is certainly not obvious and the invention described does allow effective combinations to be found in the absence of an obvious method.

- ii. Optimization for varied product applications
- iii. Robustness of approach demonstrated by similar results using different methods
- [0387] Detailed scrutiny of the results obtained by the various analyses in the body of this report, and as summarized in the tables and figures, shows the following findings.
- [0388] 1. Careful scrutiny of the performance of individual probes does not make apparent probe combinations that might perform better than any one probe alone.
- [0389] 2. All three classification methodologies evaluated hone in on similar sets of features. The small differences can be attributed to the data structure that may favor one classifier over another.
- [0390] 3. All the classifiers designed with one of these methods were shown to give good performance when tested on data from an independent pathologist, unseen during the design process. This gives high confidence in the invention.
- [0391] 4. A detection panel based on probe 7 alone gives a high performance.
- [0392] 5. If probe 7 is combined with probe 16 or 25 then a better performance is obtained.
- [0393] 6. While combinations of other probes with probe 7 appear to improve performance further, the number of extra cases captured is so low that they may be unrepresentative and the classifier so designed may not generalize.
- [0394] 7. The performance of panels selected from probes excluding probe 7 provided some discrimination, good enough in comparison with current practice using human screening, but perhaps not good enough for an automated cytometer in tomorrow's clinical diagnostic cytology world (see Figure 6).
- [0395] 8. Other combinations of probes can provide a useful, but lesser, performance.
- [0396] 9. If some probes become unavailable this invention allows the selection of other combinations of probes. This was illustrated by classifier designs based on a commercially preferred set of probes only. See Figure 7.
- [0397] 10. The invention allows a weighting to be applied against costly probes. Rather than totally excluding them from the analysis this allows their inclusion in the panel if their contribution is important.
- [0398] 11. The invention allows the design of single lung cancer type specific discrimination panels that can discriminate one type of lung cancer from among all other cancers.

[0399] 12. Analysis of the performance of a single panel to classify five cancers showed discrimination was possible but the overall error rate was worse than a set of five panels each designed to discriminate one of the cancers from the others.

- [0400] 13. A very useful discrimination was obtained with the combination of five two way classifiers.
- [0401] 14. Common sets of probes were selected by the three classification methodologies for the five discrimination panels, again giving confidence in this result.
- [0402] 15. Probes for isolating cases of Adenocarcinoma are 4, 14, 19, 20, 25, and 27.
- [0403] 16. Probes for isolating cases of Squamous Cell cancer are 1, 2, 3, 24, 25, and 26.
- [0404] 17. Probes for isolating cases of Large Cell cancer are 1 and 7, or 1 and 21.
- [0405] 18. Probes for isolating cases of Mesothelioma are 3, 12, and 16.
- [0406] 19. Probes for isolating cases of Small Cell cancer are 12, 20, and 23.
- [0407] 20. Probes for recognizing all cancers simultaneously are 1, 2, 3, 4, 12, 14, 19, 22, 23, and 28.
- [0408] 21. An advantage of using the multiple pair-wise panels as defined by this invention is that doubtful cases may not score on any of the five panels, also confusing cases may show on two or more panels. Such anomalous reports would alert the cytologist that further analysis is indicated.

iv. Risk Management Study

[0409] All the tests applied in this study were statistical in nature. There is a risk that probes selected on the basis of small improvements in performance will have statistical variations when tested on new data. To give confidence in the results, the best classifier emerging from the Linear Discriminant analysis on the Pathologist 1 and Pathologist 2 data was tested. It should be remembered that the Pathologist 3 data was statistically different from the Pathologist 1 and Pathologist 2 data, so if good performances are obtained when tests using the Pathologist 3 data, then this would be encouraging indeed.

(1) Report on Testing with unseen data - Detection panel

(a) Method

[0410] In the Section titled "Detection Panel(s) Composition" above, we showed that good classification is obtained with features 7 and 16. Using SPSS all the Pathologist 3 data that reported H scores for both 7 and 16 was selected. Then, using Transform and Compute, the

canonical discrimination function was generated as a new feature. The performance of this feature alone was then tested.

(b) Results

[0411] These are the results of testing the classifier designed on Pathologist 1 and Pathologist 2 data and testing on Pathologist 3 data. The classifier was designed using the linear discriminant function on probes 7 and 16. The Canonical Pathologist 2 function was =0.965*Probe7-0.298*Probe16.

Classification Results on Pathologist 3 data using probes 7 and 16

	٠	Total		
	Diagnosis (UCLA)	0	* 1	
Original Count	Ó	20	1	21
	_ 1	6	41	47
% .	0	95.2	4.8	100.0
	1	12.8	87.2	100.0
Cross-validated Count	0	20	1	21
	1	· . 6	41	· 47
%	0	95.2	4.8	100.0
•	1	12.8	87.2	100.0

a Cross validation is done only for those cases in the analysis. In cross validation, each case is classified by the functions derived from all cases other than that case.

[0412] This is better than classifying the Pathologist 3 data on probe 7 only show as follows:

Classification Results on Pathologist 3 data using probe 7 only

		Predicted Group	•	Total
		Membership		
•	Diagnosis (UCLA)	. 0	1	
Original Count	0	20	1	21
	1 [.]	. 8	39	47
%	0	95.2	4.8	100.0
	. 1	17.0	83.0	100.0
Cross-validated Count	0	20	1	21
	1	8	39	. 47
%	. 0	95.2	4.8	100.0
	1	17.0	83.0	100.0

a Cross validation is done only for those cases in the analysis. In cross validation, each case is classified by the functions derived from all cases other than that case.

b 89.7% of original grouped cases correctly classified.

c 89.7% of cross-validated grouped cases correctly classified.

b 86.8% of original grouped cases correctly classified.

c 86.8% of cross-validated grouped cases correctly classified.

(c) Conclusion

[0413] This gives confidence that the two-probe classifier based on 7 and 16 is better than probe 7 alone.

(2) Report on testing with unseen data – Discrimination Panel

(a) Background

[0414] Reported below is the performance of the classifier designed with Pathologist 1 and Pathologist 2 data using LDF and tested with the unseen Pathologist 3 data. The numbers of cases at the design stage was relatively small and the numbers in the test data are also small, so a good degree of variability can be expected between performance on the first and second set.

(b). Method

[0415] In SPSS, the canonical discrimination functions derived in the section titled "Pattern recognition", were built and tested on Pathologist 3 data for all five classes of cancer.

(c) Results

[0416] Mesothelioma LDF= probe3sc * .385 - probe12s * .317 + probe16s * 1.006. Classification Results

,			Predicted Group	•	Total
		Me	•		
		Meso = 1, others = 0	Ò	1	•
Original	Count	. 0	38	2 .	40
		1	1	7	8.
•	%	0.	95.0	5.0	100.0
•	•	. 1	12.5	87.5	100.0
Cross- validated	Count	. 0	. 38	2	40
		1.	-1	7	. 8
	%	0	95.0	5.0	100.0
:		1	12.5	87.5	100.0

a Cross validation is done only for those cases in the analysis. In cross validation, each case is classified by the functions derived from all cases other than that case.

b 93.8% of original grouped cases correctly classified.

c 93.8% of cross-validated grouped cases correctly classified.

[0417] Small cell cancer LDF= probe12s * .575 - probe20s * .408 - probe22s * .423 + probe23s* .344.

Classification Results

	0000	•	O !!		Total	
		Predicted				
			Group			
		Δ.	lembershi	•		
		IV	iei inei ei i		*	
			р			
		Small = 1.	0	1		
		others = 0				
Original	Count	0	39	3	42	
. •		1	1	. 5	6	
	%	. 0	92.9	7.1	100.0	
		1	16.7	83.3	100.0	
Cross-	Count	0	39	3	42	
validated						
		1.	1	. 5	6	
	%	0	92.9	7.1	100.0	
	•	1	16.7	83.3	100.0	

a Cross validation is done only for those cases in the analysis. In cross validation, each case is classified by the functions derived from all cases other than that case.

[0418] Squamous cell cancer LDF= - probe1sc * .328 - probe2sc * .295 + probe3sc * .741 + probe24s * .490 + probe25s * .393 + probe26s * .426.

Classification Results

	•		Predicted Group	·. ·	Total
		•	Membership		
	Squ	iamous =	0	. 1	
	1, 0	thers = 0			
Original	Count	0	31	4	35
		1	2	9	11
•	%	. 0	. 88.6	11.4	. 100.0
•		1	18.2	81.8	100.0
Cross-	Count	. 0	31 ·	4	35
validated					
		1	2	9	11
•	%	Ō	88.6	11.4	100.0
		1	18.2	81.8	100.0

a Cross validation is done only for those cases in the analysis. In cross validation, each case is classified by the functions derived from all cases other than that case.

b 91.7% of original grouped cases correctly classified.

c 91.7% of cross-validated grouped cases correctly classified.

b 87.0% of original grouped cases correctly classified.

c 87.0% of cross-validated grouped cases correctly classified.

[0419] Large cell cancer LDF= probelsc * .847+ probe7sc * .452.

Classification Results

Group Membership Large = 1,	14331110411011	. 1000.10				
Membership Large = 1, 0 1 others = 0 Original Count 0 23 15 1 4 5 9 0 60.5 39.5 100 1 44.4 55.6 100 Cross- Count 0 23 15 validated 1 4 5						Total
Large = 1, 0 1 others = 0 Original Count 0 23 15 1 4 5 9 0 60.5 39.5 100 1 44.4 55.6 100 Cross- Count 0 23 15 validated 1 4 5		•		Group		
others = 0 Original Count 0 23 15 1 4 5 9 0 60.5 39.5 100 1 44.4 55.6 100 Cross- Count 0 23 15 validated 1 4 5			N	1embership		
Original Count 0 23 15 1 4 5 9 0 60.5 39.5 100 1 44.4 55.6 100 Cross- Count 0 23 15 validated 1 4 5			Large = 1,	0	.1	
1 4 5 % 0 60.5 39.5 100 1 44.4 55.6 100 Cross- Count 0 23 15 validated 1 4 5			others $= 0$	٠.		• .
1 4 5 % 0 60.5 39.5 100 1 44.4 55.6 100 Cross- Count 0 23 15 validated 1 4 5	Original	Count	0	23	15	38
1 44.4 55.6 100 Cross- Count 0 23 15 validated 1 4 5			. 1	4	5	9 .
1 44.4 55.6 100 Cross- Count 0 23 15 validated 1 4 5	;	%	0	60.5	39.5	100.0
validated 1 4 5			1	44.4	55.6	100.0
1 4 5 % 0 605 395 100		Count	. 0	. 23	15	38
94 0 60.5 39.5 100			. 1	4	. 5	. 9
76 0 00.5		. %	. 0	60.5	39.5	. 100.0
1 44.4 55.6 100		•	1	44.4	55.6	100.0

a Cross validation is done only for those cases in the analysis. In cross validation, each case is classified by the functions derived from all cases other than that case.

[0420] The lower, but useful, performance was on a classifier designed and tested with a very small number of cases of large cell cancer, so this result is still very encouraging.

[0421] Adenocarcinoma, LDF= - probe4sc * .515 + probe5sc * .299 - probe14s * .485 - probe19s * .347 + probe20s * .723 + probe25s * .327 + probe27s * .327.

Classification Results

			Predicted Group Membership		Total
		Adeno = 1, Others = 0	0	1	•
Original	Count	0	29	5	34
		1.	0	14	. 14
	%	, O	·85.3	14.7	100.0
		1	0	100.0	100.0
Cross- validated	Count	0	. 29	5	34
		1	Ō	14	14
	. %	ó	85.3	14.7	100.0
		· 1	.0	100.0	100.0

a Cross validation is done only for those cases in the analysis. In cross validation, each case is classified by the functions derived from all cases other than that case.

(d) Conclusion

[0422] It is very encouraging to note the performance of these classifiers stand up to the tests of applying unseen data. This gives a very high confidence in the ability to detect the individual cancers.

b 59.6% of original grouped cases correctly classified.

c 59.6% of cross-validated grouped cases correctly classified.

b 89.6% of original grouped cases correctly classified.

c 89.6% of cross-validated grouped cases correctly classified.

(3) Training and testing on data from different patients and pathologists

[0423] As a "final final" test of robustness a LDF was trained on the data that was reviewed by both Pathologist 1 and Pathologist 2. This removes data reviewed by Pathologist 3. Hence testing on data reviewed by both Pathologist 3 plus Pathologist 1 data is not biased. Previously the test process was biased through using data from the same patient for test and train.

[0424] LDF produced the same set of features except for probe 4 which was not included. The LDF was = probe1sc * .288 + probe7sc * .846 - probe15s * .249 - probe16s * .534.

Classification Results

. .

Area under the Curve = .977

	÷		. Predicted Group		Total
		•	Membership		
		Diagnosis (UCLA)	0	1	
Original	Count	Ó	20	. 0	20 -
		1	9	37	46
•	· %	0	100.0	.0	100.0
٠.	•	1	19.6	80.4	100.0
Cross- validated	Count	0	20	0	20
	•	1	. 9	37	46
	%	. 0	100.0	.0	100.0
		1	19.6	80.4	100.0

a Cross validation is done only for those cases in the analysis. In cross validation, each case is classified by the functions derived from all cases other than that case.

b 86.4% of original grouped cases correctly classified.

c 86.4% of cross-validated grouped cases correctly classified.

[0425] Still a reasonable result, but a similar result, but with a smaller area under the curve, was obtained with probe7 alone on Pathologist 3 only data.

Classification Results

rea under the	e curve =	.908			
		F Mer	Total		
•			р		
		Diagnosis (UCLA)	0	1	
Original	Count	0	. 19	.1	20 :
•		1	. 7	39	46
	. %	.0	95.0	5.0	100.0
	. •	1	15.2	84.8	100.0
Cross- validated	Count	0	19	1	20
		1	. 7	39	46
	%	0	95.0	5.0	100.0
		1	15.2	84.8	100.0

a Cross validation is done only for those cases in the analysis. In cross validation, each case is classified by the functions derived from all cases other than that case.

II. Colorectal Cancer

[0426] Epithelial tumors of intestines are a major cause of morbidity and mortality worldwide. The colon (including the rectum) is host to more primary neoplasms than any other organ in the body. Colorectal cancer ranks second only to bronchogenic carcinoma among the cancer killers. Adenocarcinomas constitute the vast majority of colorectal cancers and represent 70% of all malignancies arising in the gastrointestinal (GI) tract. The small intestine is an uncommon site for benign or malignant tumors despite its great length and vast pool of dividing mucosa cells. (Crawford, J.M., The Gastrointestinal Tract, in Robbins Pathologic Basis of Disease, R.S.e.a. Cotran, Editor. 1999, W. B. Saunders Company: Philadelphia. p. 775-843).

[0427] The peak incidence for colorectal carcinomas is in the patient age range of 60 to 79 years. Fewer than 20% of cases occur before the age of 50 years. When colorectal carcinoma is found in a young person, pre-existing ulcerative colitis or one of the polyposis syndromes must be suspected. Colorectal carcinoma has worldwide distribution. The highest death rates are found in the United States and Eastern European countries, up to 10-fold greater than the rates in Mexico, South America and Africa. Environmental factors, particularly dietary practices, are implicated in these striking geographic contrasts. In addition, many studies implicate obesity and physical inactivity as risk factors for colon cancer. (Crawford, J.M., *The Gastrointestinal Tract*,

b 87.9% of original grouped cases correctly classified.

c 87.9% of cross-validated grouped cases correctly classified.

in Robbins Pathologic Basis of Disease, R.S.E.A. Cotran, Editor. 1999, W. B. Saunders Company: Philadelphia. p. 775-843).

[0428] Almost all cancers (98%) found in the large intestine are adenocarcinomas. Virtually all colorectal carcinomas exhibit genetic alterations, including E-cadherin and β-catenin in adenomatous polyposis coli (APC); human mismatch repair genes, hMSH2, hMLH1 and hPMS2 in hereditary non-polyposis colon carcinoma (HNPCC); and, mutation of K-ras and p53 genes. (Crawford, J.M., The Gastrointestinal Tract, in Robbins Pathologic Basis of Disease, R.S.E.A. Cotran, Editor. 1999, W. B. Saunders Company: Philadelphia. p. 775-843). Diagnosing colorectal carcinoma in an early stage is one of the prime challenges to medical professionals because these carcinomas present with unspecific clinical symptoms such as fatigue, anemia, abdominal pain and bloody stools. Currently, the main diagnostic method is colonoscopy to visually examine whether there is a tumor mass. However, this is an invasive procedure that involves colon-prepping by the patient and anesthesia throughout the procedure to achieve unconsciousness. It is not surprising that patient compliance is a major issue. Unfortunately, available fecal occult blood tests or the Guaic tests are not specific enough. Therefore, current detection methods for colorectal cancer, such as colonoscopy or sigmoidoscopy, have proven to be inadequate screening tools due to the invasiveness of the procedures, the relative lack of accuracy and poor patient compliance. Furthermore, non-invasive fecal occult blood testing (FOBT) is not effective and suffers from lack of sensitivity or specificity.

[0429] Molecular diagnosis of colorectal carcinoma receives much attention because most of these cancers have genetic abnormalities. Among technologies, immunohistochemistry (IHC) and immunocytochemistry (ICC) are widely used to evaluate colorectal carcinogenesis. A variety of colorectal tumor markers have been discovered to aid physicians in making timely, precise diagnoses, and to provide significantly better patient management. Unfortunately, none of these tumor markers is a "magic bullet" with both a high sensitivity and specificity. Therefore, alternative ways to enhance diagnostic accuracy are necessary.

[0430] An alternative way to enhance diagnostic accuracy is to develop a panel comprising a plurality of probes each of which specifically binds a marker associate with colorectal cancer. All candidate probes are to be tested with ICC techniques. In some embodiments, specimens may be obtained from colonic washings. Although cytological specimens obtained from colonic washings often have fewer cells than tissue sections, the use of high quality polyclonal or

monoclonal antibodies may be employed to ensure good assay performance. In some embodiments, test slides may be made by spiking tumor cells into a cell suspension before actual patient specimens are tested. In other embodiments, limitations due to colonic washings often having fewer cells than tissues sections may be overcome by studying patients who match the variables as closely as possible, such as age, gender, diagnosis, tumor grade, tumor size, clinical stage, etc.

Statistical analysis will be used to design panels, as described above for lung cancer. During processing, technical issues such as cell smears or pellets not sticking to slides during harsh washings may occur in some embodiments. However, such issues can readily be addressed by manipulation of software or modifying staining protocols to mitigate such problems. In some embodiments, the specimens will be processed and analyzed using a device that automatically samples the specimen and prepares slides for diagnosis. It is anticipated that a broad menu of probes will be used initially. The number of probes will be pruned to a suitably sized panel in order to retain a high level of sensitivity and specificity. Selection of the final probes will be based on a pre-defined threshold of the percentage of positive stained tumor cells. Sophisticated statistical analysis will be employed to make these determinations. Since the panel-assay approach to detecting malignancies is applicable to solid tumors, and several of the same tumor markers are in different panels, this method may be carried out in parallel, as well as serially. In this manner, the assay development process can be expedited.

[0432] Compared with lung cancers, which have five subtypes (adenocarcinoma, squamous cell carcinoma, small cell carcinoma, large cell carcinoma, and mesothelioma), colorectal epithelial tumors are predominantly adenocarcinoma. This allows the colorectal tumor panel to be specifically targeted at only one type of cancer. A large number of cytological specimens is not necessary because the panel can be tested on either biopsied or colectomy tissue.

Library of Probes/Markers

[0433] Various sources containing information about cancer markers were reviewed. An arbitrary criterion of 20% or greater positivity of colorectal carcinoma was used to select probes for a preferred panel for detection and/or diagnosis of colorectal cancer. The term "20% or greater positivity" means that if 100 tumor cases were studied, 20 or more of these cases would have shown a presence of the individual marker, while the remaining 80 cases would not have

shown a presence of the individual marker. A preferred panel may include molecular markers selected from AKT, β-catenin, Brain-type Glycogen Phopshorylase (BPG), Caveolin -1, CD44v6, cFLIP, Cripto-I, Amphiregulin, Cyclin D1, Cyclooxygenase (COX-2), Cytokeratin 20 (CK20), Carcinoembryonic Antigen (CEA), E-cadherin, Bcl-2, Bax, hMLH1, hMSH2, Epidermal Growth Factor Receptor (EGFR), Ephrin-B2 (Eph-B2), Ephrin-B4 (Eph-B4), FasL, HMGI(Y), Ki-67, Lysozyme, Matrilysin (MMP-7), p16, p68, Retinoblastoma (Rb), cdk2/cdc2, S100A4, YB-1 and p53. A brief description of the library of probes/markers utilized in the present example is provided below.

AKT

[0434] This is a proto-oncogen with profound anti-apoptotic activity. This serine-threonine kinase is over-expressed in numerous malignancies. Roy et al showed normal colonic mucosa and hyperplastic polyps exhibited no significant AKT expression, in marked contrast to the dramatic AKT immunoreactivity seen in colorectal cancers (57% positive). In addition, AKT was also detected in 57% of adenomas indicating over-expression of this proto-oncogen as an early event during colon carcinogenesis. (Roy, H.K., et al., AKT proto-oncogene overexpression is an early event during sporadic colon carcinogenesis. Carcinogenesis, 2002. 23(1): p. 201-5).

B-catenin

[0435] Tumor-suppressor gene, adenomatous polyposis coli (APC), has been detected in 80% of sporadic colorectal carcinomas. They occur in small-sized adenomas and even in the smallest lesions with the risk of neoplasia, the dysplastic aberrant crypt foci. Products of the APC gene regulate intracellular β -catenin. Studies have shown that mutated APC causes decreased turnover rate and leads to an accumulation of β -catenin. Herter et al discovered not only a linear increased expression level of β -catenin but also a different location of β -catenin from adenoma to carcinoma, in a majority of cases. In adenomas with mild and moderate dysplasia, β -catenin is only present in the nucleus. In severe dysplastic adenomas and carcinomas, it is present in both cytoplasm and the nucleus. In normal colonic mucosa, the only weak staining is in the cell-to-cell border membranes and cytoplasm. These results may be important for diagnostic and clinical purpose, because the nuclear presence of β -catenin may be the earliest molecular evidence of colorectal malignancy. (Herter, P., et al., Intracellular distribution of beta-catenin in colorectal adenomas, carcinomas and Peutz-Jeghers polyps. J Cancer Res Clin Oncol, 1999. 125(5): p. 297-304).

Brain-type Glycogen Phosphorylase (BPG)

[0436] Brain-type glycogen phosphorylase (BPG) is a unique intestinal malignancy marker. It has three major isoforms: muscle, liver and brain. Previously, gastric carcinoma has been shown to be associated with abnormal expression of BPG through immuno-histochemistry (IHC) techniques. A new study by Tashima et al demonstrated for the first time that BPG was present in 83% of colorectal carcinoma by IHC. And more interestingly, normal colonic mucosa from remote sites are negative for BPG whereas overtly normal-looking mucosa adjacent to a tumor are positive. (Tashima, S., et al., Expression of brain-type glycogen phosphorylase is a potentially novel early biomarker in the carcinogenesis of human colorectal carcinomas. Am J Gastroenterol, 2000. 95(1): p. 255-63).

Caveolin-1

[0437] This is a major structural protein of caveolae, the vesicular invaginations of the plasma membrane. Studies have demonstrated that caveolin family members contain a common domain, termed caveolin-scaffolding, that functions to organize signaling molecules, including G-protein, Ha-ras, Src-family tyrosin kinases, and epidermal growth factor receptor (EGFR). While in vitro and in vivo animal experiments demonstrated a suppressive effect of caveolin-1 in cell transformation and breast carcinogenesis, other studies, including studies of human breast and prostate cancers, revealed a positive association of caveolin-1 expression with tumorigenesis and progression, suggesting a tumor-promoting function. Fine et al's study showed 88% of colonic adenocarcinomas are positive for caveolin-1 by IHC. (Fine, S.W., et al., Elevated expression of caveolin-1 in adenocarcinoma of the colon. Am J Clin Pathol, 2001. 115(5): p. 719-24).

CD44v6

[0438] This is a widely expressed cell-surface glycoprotein that may be involved in cell-to-cell and cell-to-matrix interactions. An abundance of CD44s is present on cells of normal epithelial and hematopoietic origin. In contrast, the alternatively spliced CD44 variants (CD44v) are expressed predominantly on cells and tumors of epithelial origin. Several reports have shown expression of CD44v6 with an advanced stage of colorectal carcinoma. Ishida studied 63 colorectal carcinoma patients through IHC techniques and found 59% of the cases to be positive for CD44v6. Normal colonic mucosa are negative for CD44v6. (Ishida, T., Immunohistochemical expression of the CD44 variant 6 in colorectal adenocarcinoma. Surg Today, 2000. 30(1): p. 28-32).

Cellular FLICE-like Inhibitory Protein (cFLIP)

[0439] Cellular FLICE-like inhibitory protein (cFLIP) is an endogenous inhibitory regulator of Fas-mediated apoptosis. Although the physiological functions of cFLIP have not yet been clarified, pathogenetic implications in disease, including tumors, have been suspected. Moreover, it has recently been reported that overexpression of cFLIP results in the escape of tumor cells from T-cell immunity and is possibly related to tumor establishment and growth. Ryu et al have demonstrated that 100% (52/52) of colonic adenocarcinomas are positive for cFLIP. Normal colonic epithelium, adeonomatous polyps have much lower staining intensity. (Ryu, B.K., et al., *Increased expression of cFLIP(L) in colonic adenocarcinoma*. J Pathol, 2001. 194(1): p. 15-9).

Cripto-I and Amphiregulin

[0440] Cripto-I (CR-I) and amphiregulin (AR) are epidermal growth factor (EGF)-related peptides. Several reports have demonstrated that AR and CR-I function as autocrine growth factors in human colon epithelial cells in vitro. Furthermore, it has been demonstrated that AR and CR-I are expressed in a majority of human primary colon carcinomas. In particular, overexpression of either AR or CR-I proteins has been found by IHC in approximately 70% of human colon adenomas and carcinomas. (De Angelis, E., et al., Expression of cripto and amphiregulin in colon mucosa from high risk colon cancer families. Int J Oncol, 1999. 14(3): p. 437-40).

Cyclin D1

[0441] This protein plays an important role in cell proliferation. Mutations and/or altered expression of Cyclin D1 are involved in neoplasia. Increased expression of Cyclin D1 is observed in esophageal, head, neck, hepatic, breast and colorectal cancers. A study by Arber et al revealed increased Cyclin D1 staining in 30% of colorectal adenocarcinomas and 34% of adenomatous polyps but not in hyperplastic polyps or normal mucosa. (Arber, N., et al., Increased expression of cyclin D1 is an early event in multistage colorectal carcinogenesis. Gastroenterology, 1996. 110(3): p. 669-74).

Cyclooxygenase (COX-2)

[0442] This is a prostaglandin synthase enzyme involved in arachidonic acid metabolism.

Since evidence showed tumor-suppressive effects of nonsteroidal antiinflammatory drugs

(NSAIDs) on colorectal cancer, Cyclooxygenase has received attention because it is plausible

that the tumor-suppressive effects of NSAIDs are due to their reduction of COX-2 activity. Sakuma et al demonstrated that 38% of colorectal cancers are COX-2 positive. COX-2 staining was more intense in areas of cancer tissue than in non-cancerous tissues. In some specimens, tissue close to the cancer was stained more intensively than tissue further away from the cancer. (Sakuma, K., et al., Cyclooxygenase (COX)-2 immunoreactivity and relationship to p53 and Ki-67 expression in colorectal cancer. J Gastroenterol, 1999. 34(2): p. 189-94).

Cytokeratin 20 (CK 20) and Carcinoembryonic antigen (CEA)

[0443] These are well known colorectal carcinoma markers. They have been reported in many scientific articles. Even though they are not specific for colorectal tumor, one study by Lagendijk et al revealed that immunohistochemically CK 20 and CEA are two key discriminative markers to differentiate metastatic colonic adenocarcinoma to ovary from breast and primary ovarian carcinoma. (Lagendijk, J.H., et al., Immunohistochemical differentiation between primary adenocarcinomas of the ovary and ovarian metastases of colonic and breast origin.

Comparison between a statistical and an intuitive approach. J Clin Pathol, 1999. 52(4): p. 283-90).

E-cadherin, p53, Bcl-2, Bax, hMLH1, hMSH2

[0444] Kapiteijn et al and Bukholm et al, in separate studies discovered a number of oncogenes and tumor suppressor genes involved in the oncogenesis of colorectal cancers. E-cadherin, p53, Bcl-2, Bax, all showed greater than 20% immunostaining in tumor cells. Mismatch repair genes, hMLH1 and hMSH2 are also significantly increased. These repair genes are involved in genetic "proof-reading" during DNA replication, and hence are referred to as caretaker genes. Mutation of these genes has been shown to be involved in the early development of gastrointestinal malignancy. (Kapiteijn, E., et al., Mechanisms of oncogenesis in colon versus rectal cancer. J Pathol, 2001. 195(2): p. 171-8; Bukholm, I.K. and J.M. Nesland, Protein expression of p53, p21 (WAF1/CIP1), bcl-2, Bax, cyclin D1 and pRb in human colon carcinomas. Virchows Arch, 2000. 436(3): p. 224-8). Yantiss et al showed p53 and E-cadherin are two markers differentiating adenomas with misplaced epithelium from adenomas with invasive adeocarcinoma, indicating the specificity of these two markers for colorectal cells. (Yantiss, R.K., et al., Utility of MMP-1, p53, E-cadherin, and collagen IV immunohistochemical stains in the differential diagnosis of adenomas with misplaced epithelium versus adenomas with

invasive adenocarcinoma. Am J Surg Pathol, 2002. 26(2): p. 206-15). These markers are common to almost all the solid tumors.

Epidermal Growth Factor Receptor (EGFR)

[0445] Epidermal growth factor receptor (EGFR) is a 170-kilodalton transmembrane cell-surface receptor. It, along with c-erb B-2, c-erb B3, and c-erb B4, has tyrosine kinase activity and is encoded by the c-erb-B protooncogene. Chimeric anti-EGFR monoclonal antibody is an investigational therapy for advanced stages of colon adenocarcinoma. Increased levels of EGFR are found in many solid tumors, including colorectal carcinoma, squamous cell carcinoma of the lung, head, neck, cervix, breast, prostate and bladder. One study by Goldstein et al showed 75% of the colonic adenocarcinoma had EGFR immunohistochemical positivity (Goldstein, N.S. and M. Armin, Epidermal growth factor receptor immunohistochemical reactivity in patients with American Joint Committee on Cancer Stage IV colon adenocarcinoma: implications for a standardized scoring system. Cancer, 2001. 92(5): p. 1331-46).

Ephrin-B2 (Eph-B2) and Ephrin-B4 (Eph-B4)

[0446] The erythropoietin-producing amplified sequence (Eph) family is the largest sub-family of receptor tyrosine kinases (RTKs). Eph-B2 and B4 are the ligands binding to Eph. The ephrin-Eph system is important in embryological development and differentiation of the nervous and vascular systems. Several studies have shown that high expression of ephrins may be associated with increased potential for tumor growth, tumorigenicity, and metastasis. Using immunohistochemical analysis, Liu et al showed Eph-B2 and Eph-B4 had greater staining intensity in 100% (5/5) of the cases studied compared with adjacent normal mucosa. (Liu, W., et al., Coexpression of ephrin-Bs and their receptors in colon carcinoma. Cancer, 2002. 94(4): p. 934-9).

FasL

[0447] This is a transmembrane protein member of the tumor necrosis factor super-family, and induces cell death in apoptosis-sensitive cells expressing its receptor, Fas (CD95/ APO-I). It has been widely demonstrated that FasL is up-regulated in several types of cancer. Moreover, in vitro and in vivo studies have shown that FasL can enable cancer cells to mount a Fas counterattack, impairing the immune response by inducing apoptosis in anti-tumor immune effector cells. These findings suggest that FasL expression by cancer cells may be an important factor in the inhibition of anti-tumor immune responses. Belluco et al showed that FasL

expression is a relatively early event of carcinomas and in colorectal tumorigenesis. FasL expression was found in 28% of hyperplastic polyps, 76% of low grades and 93% of high-grade polyps (Belluco, C., et al., Fas ligand is up-regulated during the colorectal adenoma-carcinoma sequence. Eur J Surg Oncol, 2002. 28(2): p. 120-5). The results are in line with others findings that FasL expression was detected in 81% of carcinomas and in 41% of adenomas. Moreover, FasL was significantly more frequently expressed in high-grade dysplastic adenomas than in low-grade adenomas.

HMGI(Y)

[0448] Proteins HMG-I, HMG-Y and HMGI-C constitute the high mobility group I protein family. The first two proteins are encoded by the same gene, HMGI(Y), through alternative splicing, while HMGI-C is the product of a different gene. HMGI genes are involved in the generation of benign and malignant tumors. Previous reports showed HMGI(Y) proteins are abundantly expressed in colon carcinoma cell lines and tissues but not in normal colon mucosa. Chiappetta et al discovered 36 colorectal carcinomas were all positive for HMGI(Y) by IHC, whereas no expression was detected in normal colon mucosa. HMGI(Y) expression in adenomas was closely correlated with the degree of cellular atypia. Only 2 of the 18 non-neoplastic polyps tested were HMGI(Y)-positive. (Chiappetta, G., et al., High mobility group HMGI(Y) protein expression in human colorectal hyperplastic and neoplastic diseases. Int J Cancer, 2001. 91(2): p. 147-51). These results indicate that HMGI(Y) protein induction is associated with the early stages of neoplastic transformation of colon cells and only rarely with colon cell hyperproliferation.

Ki-67'

[0449] Ki-67 is a cell proliferation nuclear marker. It is expressed in a variety of tumors, including colorectal cancer. One study by Sakuma et al demonstrated 48% of colorectal cancers are positive for Ki-67. (Sakuma, K., et al., Cyclooxygenase (COX)-2 immunoreactivity and relationship to p53 and Ki-67 expression in colorectal cancer. J Gastroenterol, 1999. 34(2): p. 189-94).

Lysozyme

[0450] Lysozyme is an enzyme with a broad spectrum of antibacterial activities. It is present in numerous human tissue fluids and secretions, including saliva, tears, mils, serum, and gastric and small intestinal juice. Lysozyme is absent in normal colonic epithelium. Interestingly, numerous

immunohistochemical studies demonstrated the expression of lysozyme in the tumor cells of gastric adenomas and adenocarcinomas. Many studies have shown lysozyme positivity in colon cancer ranging from 28% to 80% while normal colonic glands adjacent to the adenocarcinomas did not show any lysozyme protein expression. (Yuen, S.T., et al., *Up-regulation of lysozyme production in colonic adenomas and adenocarcinomas*. Histopathology, 1998. 32(2): p. 126-32).

Matrilysin (MMP-7)

[0451] Matrilysin is a member of the MMP gene family and has proteolytic activity against a spectrum of substrates, such as collagens, proteoglycans, elastin, laminin, fibronectin and casein. It is produced by malignant tumor cells such as esophageal, colorectal, gastric, head, neck, lung, prostate and heptocellular carcinomas. Immunohisto-chemical studies have shown that the expression of matrilysin correlates significantly with nodal or distant metastasis in gastric and colorectal carcinomas. A study by Masaki et al showed 34% of colorectal carcinomas are positive for matrilysin. (Masaki, T., et al., Matrilysin (MMP-7) as a significant determinant of malignant potential of early invasive colorectal carcinomas. Br J Cancer, 2001. 84(10): p. 1317-21).

p16

[0452] This is a cell cycle inhibitor and a major tumor-suppressor protein. A role for p16 in intestinal neoplasia is suggested by the observation that the promoter region is methylated in a subset of human colon tumors. Dai et al showed p16 expression was very low in normal mucosa, and in 18 of 28 primary colon carcinomas and 5 of 5 metastatic colon carcinomas. In addition, p16 staining correlated inversely with that of Ki-67, cyclin A and the retinoblastoma protein, suggesting cell cycle progression was inhibited. (Dai, C.Y., et al., p16(INK4a) expression begins early in human colon neoplasia and correlates inversely with markers of cell proliferation.

Gastroenterology, 2000. 119(4): p. 929-42).

p68

[0453] This is an interferon-inducible protein kinase, which is a key factor in the regulation of both viral and cellular protein synthesis. Its expression is correlated with cellular differentiation in both normal and neoplastic cell types. Singh et al have found that p68 is positive in 76% of colorectal carcinoma patients. Normal colonic mucosa showed weak p68 staining. High p68 expression demonstrated a trend toward improved survival. Patients with tumors expressing high levels (3 to 4+) of p68 had a longer 5-year survival rate compared to patients with lower p68

expression. (Singh, C., et al., Expression of p68 in human colon cancer. Tumour Biol, 1995. 16(5): p. 281-9).

Rb and cdk2/cdc2

[0454] The retinoblastoma (Rb) gene is a tumor-suppressor gene and its product, pRB, is known to act as a negative regulator of the cell cycle. Although lack of pRB expression resulting from gene alterations is considered to be responsible for the genesis of several malignancies, including osteosarcomas and carcinomas of the lung, breast and bladder. In contrast, colorectal cancer has reportedly shown infrequent inactivation of this gene, and Southern blot analysis has demonstrated Rb gene amplification in approximately 30% of colorectal cancers. Yamamoto et all studied Rb and its related kinases, cdk2 and cdc2, by Western blot and found increased levels of cdk2/cdc2, as well as hyperphosporylated form of pRB in colorectal carcinoma. Furthermore, immunohistochemical studies showed that cdc2/cdc2 was expressed exclusively in cancer cells positive for pRB. These results suggest that an increase in the expression of cdk2/cdc2 in colorectal cancer may have prevented pRB from braking the cell cycle through phosphorylation. (Yamamoto, H., et al., Coexpression of cdk2/cdc2 and retinoblastoma gene products in colorectal cancer. Br J Cancer, 1995. 71(6): p. 1231-6).

S100A4

[0455] This is a calcium-binding protein and has been implied to be involved in cell immortalization, cell growth, differentiation of mammary epithelial stem cells to myoepithelial-like cells, and fibrogenesis. In addition, S100A4 has been reported to be specifically expressed in metastatic tumor cells. Takenaga et all observed 44% of focal carcinomas and 94% of adenocarcinomas were immunopositive while none of the adenomas were positive. Interestingly, the incidence of immunopositive cells increased according to the depth of invasion, and nearly all of the carcinoma cells in 14 metastases of the liver were positive. These results suggest that S100A4 is a good marker to differentiate adenoma from adenocarcinoma and may be involved in the progression and metastatic process of colorectal neoplastic cells. (Takenaga, K., et al., Increased expression of S100A4, a metastasis-associated gene, in human colorectal adenocarcinomas. Clin Cancer Res., 1997, 3(12 Pt 1): p. 2309-16).

Y-box Binding Protein (YB-1)

[0456] The Y-box binding protein (YB-1) is a member of a family of DNA binding proteins that contain a highly conserved, cold shock domain and interacts with inverted CCAAT boxes (Y

boxes). YB-1 is expressed in a wide range of cell types and has been implicated in the regulation of various genes involved in cell proliferation. It is also overexpressed in cisplatin-resistant cancer cell lines, suggesting that YB-1 may be involved in either DNA repair or DNA damage response, in addition to its role as a transcription factor. Shibao et al showed YB-1 was overexpressed in almost all cases of colorectal carcinomas compared with normal mucosa. (Shibao, K., et al., Enhanced coexpression of YB-1 and DNA topoisomerase II alpha genes in human colorectal carcinomas. Int J Cancer, 1999. 83(6): p. 732-7).

Utility of Colorectal Carcinoma Panel

[0457] The colorectal carcinoma panel contains not only markers for early detection of colorectal carcinoma but also markers for assessing metastatic potential and prognosis. For example, positive CEA (carcinoembryonic antigen) reaction has a significant relationship with the grade of differentiation of colorectal carcinoma while diffuse cellular expression of this antigen often indicates neoplasms extending beyond the intestinal wall and invading the lymph vessels. The number of tissue antigens expressed is significantly related to the extent of tumor spread through the intestinal wall (Lorenzi, M., et al., Histopathological and prognostic evaluation of immunohistochemical findings in colorectal cancer. Int J Biol Markers, 1997. 12(2): p. 68-74). Also, serum CEA levels and the expression of p53 proteins provide complementary prognostic information for colorectal cancer. Positive immunostaining of p53 and elevated CEA levels are associated with low cumulative disease free survival and have been shown to have independent prognostic significance (Diez, M., et al., Time-dependency of the prognostic effect of carcinoembryonic antigen and p53 protein in colorectal adenocarcinoma. Cancer, 2000. 88(1): p. 35-41). Nasierowska-Guttmejer and associates (Nasierowska-Guttmejer, A., The comparison of immunohistochemical proliferation and apoptosis markers in rectal carcinoma treated surgically or by preoperative radio-chemotherapy. Pol J Pathol, 2001. 52(1-2): p. 53-61) have shown that low expression of Ki-67 and high levels of Bax expression are correlated with the total, or near-total, response of colorectal cancer to the treatment and regression of the tumor mass. However, less than two-thirds of the cases are correlated with low expression of p53, MIB1, bax and bcl-2. Another study showed that higher p53 and Ki67 values were associated with prognostically poor histopathologic features (Saleh, H.A., H. Jackson, and M. Banerjee, Immunohistochemical expression of bcl-2 and p53 oncoproteins: correlation with Ki67 proliferation index and prognostic histopathologic parameters in colorectal neoplasia.

Appl Immunohistochem Mol Morphol, 2000. 8(3): p. 175-82). Additionally, expression of cyclin D1, CD44v6, and matrilysin (MMP-7) in colorectal cancer has been shown to be correlated with high recurrent rates, reduced relapse-free and overall poor survival (McKay, J.A., et al., Analysis of key cell-cycle checkpoint proteins in colorectal tumours. J Pathol, 2002. 196(4): p. 386-93; Ropponen, K.M., et al., Expression of CD44 and variant proteins in human colorectal cancer and its relevance for prognosis. Scand J Gastroenterol, 1998. 33(3): p. 301-9; Bhataydekar, J.M., et al., Molecular markers are predictors of recurrence and survival in patients with Dukes B and Dukes C colorectal adenocarcinoma. Dis Colon Rectum, 2001. 44(4): p. 523-3; and Adachi, Y., et al., Clinicopathologic and prognostic significance of matrilysin expression at the invasive front in human colorectal cancers. Int J Cancer, 2001. 95(5): p. 290-4). Studies also demonstrated that increased expression of p53, MMP-7, \u03b3-catenin and reduced expression pf E-Cadherin in colorectal carcinomas were associated with an increase in their metastatic potential (Zeng. Z.S., et al., Matrix metalloproteinase-7 expression in colorectal cancer liver metastases: evidence for involvement of MMP-7 activation in human cancer metastases. Clin Cancer Res, 2002. 8(1): p. 144-8; Ikeguchi, M., et al., Reduced E-cadherin expression and enlargement of cancer nuclei strongly correlate with hematogenic metastasis in colorectal adenocarcinoma. Scand J Gastroenterol, 2000. 35(8): p. 839-46; Sory, A., et al., Does p53 overexpression cause metastases in early invasive colorectal adenocarcinoma? Eur J Surg, 1997. 163(9): p. 685-92; and Hiscox, S. and W.G. Jiang, Expression of E-cadherin, alpha, beta and gamma-catenin in human colorectal cancer. Anticancer Res, 1997. 17(2B): p. 1349-54). Thus, colorectal panels will have diagnostic and prognostic value and provide patient risk stratification to guide in clinical therapy.

Preferred Probes/Markers

[0458] A preferred panel for detecting and/or diagnosis colorectal carcinoma comprises one or more tumor markers listed above. A more preferred panel for detecting and/or diagnosing colorectal carcinoma comprises one or more tumor markers selected from β-catenin, E-cadherin, hMSH2, hMLH1, p53, and cytokeratin 20. Virtually all of the colorectal carcinomas exhibit genetic alterations, thus providing an opportunity for early detection. E-Cadherin and β-catenin are intimately interacting with the APC (adenomatous polyposis Coli) tumor-suppressor gene. A defect in the APC gene is associated with FAP (familial adenomatous polyposis) and Gardner syndrome, which have very high incidence of colorectal cancer. Another genetic alteration in

colorectal carcinoma involves the DNA repair genes. Two important mismatch repair genes, hMSH2 and hMLH1, are responsible for "proof-reading" during DNA replication. Another marker, p53, is located at chromosome 17 and more than 70% of colorectal cancers have losses at chromosome 17p. Cytokeratin 20 (CK20) is consistently expressed in colorectal carcinoma and when coupled with negative cytokeratin 7 (CK7) staining, the combination (CK20+/CK7-) is highly specific for colorectal carcinoma (Chu, P., E. Wu, and L.M. Weiss, Cytokeratin 7 and cytokeratin 20 expression in epithelial neoplasms: a survey of 435 cases. Mod Pathol, 2000. 13(9): p. 962-72). The importance of these tumor markers with respect to colorectal carcinoma is detailed below.

β-catenin

[0459] The inherited defect underlying familial adenomatous polyposis (FAP) and Gardner syndromes has been mapped to 5q21, site of the APC tumor-suppressor gene. APC protein binds to cytoskeletal protein β-catenin in a cellular adhesion molecular complex, which includes intercellular adhesion molecule E-cadherin. β-catenin can also act as an oncogene. When it is not bound to E-cadherin (thus participating in cell-cell adhesion), β-catenin binds to a t-family of protein partners known as T cell factor-lymphoid enhancer factor (Tcf-Lef) proteins, which activate other genes. Genes activated by this β-catenin:Tcf complex are thought to include those stimulating cell proliferation and inhibiting apoptosis. APC binding to β-catenin directs toward degradation, thereby inhibiting the β-catenin:Tcf signaling pathway. Mutations in the APC gene reduce the affinity of APC protein for β-catenin, leading to loss of intercellular contact on the one hand and an increased cytoplasmic pool of β-catenin on the other. The resultant enhancement of Tcf-mediated cell proliferation initiates a sequence of events that predisposes to the development of carcinoma (Peifer, M., Beta-catenin as oncogene: the smoking gun. Science, 1997. 275(5307): p. 1752-3). Hence, APC is regarded as a "gatekeeper" gene. Mutations in APC underlie FAP, and are early events in the evolution of sporadic colon cancer, with mutations being found in 85% of colorectal carcinomas (Crawford, J.M., The Gastrointestinal Tract, in Robbins Pathologic Basis of Disease, R.S.e.a. Cotran, Editor. 1999, W. B. Saunders Company: Philadelphia, p. 775-843). Notably, most of the tumors without mutations in APC show mutations in β-catenin.

hMSH2 and hMLH1

[0460] Inherited mutations in any of the genes that are involved in DNA repair are putatively responsible for the familial syndrome of HNPCC. These mismatch repair genes, hMSH2, hMLH1 are involved in genetic "proof-reading" during DNA replication, and hence are referred to as "caretaker" genes. There are 50,000 to 100,000 dinucleotide repeat sequences in the human genome, and mutations in mismatch repair genes can be detected by the presence of widespread alterations in these repeats. This is referred to as microsatellite instability. Patients who inherit a mutant DNA repair gene have normal repair activity because of the remaining normal allele. However, cells in some organs (colon, stomach, endometrium) are susceptible to a second, somatic mutation that inactivates the wild-type allele. Mutation rates up to 1000 times normal ensue, such that most of the HNPCC tumors show microsatellite instability. Mutation of these genes has been shown to be involved in the early development of gastrointestinal cancer (Kapiteijn, E., et al., Mechanisms of oncogenesis in colon versus rectal cancer. J Pathol, 2001. 195(2): p. 171-8; Bukholm, I.K. and J.M. Nesland, Protein expression of p53, p21 (WAF1/CIP1), bcl-2, Bax, cyclin D1 and pRb in human colon carcinomas. Virchows Arch, 2000. 436(3): p. 224-8).

p53

[0461] Losses at chromosome 17p have been found in 70 to 80% of colon cancers. These chromosomal deletions affect the p53 gene, suggesting that mutations in p53 occur late in colon carcinogenesis. Also well known, p53 plays a critical role in cell cycle regulation. A multi-hit concept for colon cancer carcinogenesis is hypothesized. APC mutations are usually the earliest and possibly the initiating event in about 80% of sporadic colon cancers, with a less frequent contribution from mutations in mis-match repair genes. During the ensuing progression from adenoma to carcinoma, additional mutations ensue, such as late mutations or loss of heterozygosity (LOH) at p53 on chromosome 17p and in the DCC region on chromosome 18q. Cumulative alterations in the genome thus lead to progressive increases in size, level of dysplasia, and invasive potential of neoplastic lesions (Crawford, J.M., *The Gastrointestinal Tract*, in *Robbins Pathologic Basis of Disease*, R.S.e.a. Cotran, Editor. 1999, W. B. Saunders Company: Philadelphia. p. 775-843).

Cytokeratin 20 (CK 20)

[0462] CK 20 is a low molecular weight, intermediate filament. It is of particular interest because of its restricted range of expression. CK 20 is consistently expressed in normal and malignant epithelia. Expression is restricted to the gastric and intestinal epithelium and Merkel cells. Studies surveying hundreds of epithelial neoplasms from various organ systems by immunohistochemistry techniques demonstrated that virtually all cases of colorectal carcinomas are CK 20 positive (Chu, P., E. Wu, and L.M. Weiss, Cytokeratin 7 and cytokeratin 20 expression in epithelial neoplasms: a survey of 435 cases. Mod Pathol, 2000. 13(9): p. 962-72). The combination of CK 20/CK 7 immunoprofile is particularly useful in identifying the primary site of metastatic tumor in cytologic specimens. CK 20+/CK 7- is only observed in cell blocks in which colorectal was the primary site (Blumenfeld, W., et al., Utility of cytokeratin 7 and 20 subset analysis as an aid in the identification of primary site of origin of malignancy in cytologic specimens. Diagn Cytopathol, 1999. 20(2): p. 63-6). Ascoli and associates (Ascoli, V., et al., Utility of cytokeratin 20 in identifying the origin of metastatic carcinomas in effusions. Diagn Cytopathol, 1995. 12(4): p. 303-8) determined that CK 20 expression by immunohistochemistry was consistently seen in malignant effusions from colonic origin.

[0463] The following abbreviations were used throughout this example: APC = Adenomatous Polyposis Coli, AR = Amphiregulin, BGP = Brain-type glycogen phosphorylase, CD44v6 = CD44 Splice Variant 6, CEA = Carcinoembryonic Antigen, CFLIP = Cellular FLICE-like inhibitory protein, CK = Cytokeratin, COX = Cyclooxygenase, CR-I = Cripto, EGFR = Epidermal Growth Factor Receptor, EphB = Erythropoietin-Producing Amplified Sequence, FasL = Fas Ligand, FOBT = Fecal Occult Blood Test, HNPCC = Hereditary Nonpolyposis Colon Cancer, ICC = Immunocytochemistry, IHC = Immunohistochemistry, MLH = Human Mismatch Repair Genes, MMP = Matrilysin (MMP-7), MSH = Human Mismatch Repair Genes, PLAP = Placental Alkaline Phosphatase, Rb = Retinoblastoma and YB = Y-box Binding Protein.

III. Bladder Cancer

[0464] Neoplasms of the bladder pose biological and clinical challenges. The incidence of these epithelial tumors in the United States has been steadily increasing during the past few years and now amounts to more than 50,000 new cases annually. Despite improvements in detection and management of these neoplasms, the death toll remains at about 10,000 annually (Crawford, J.M. and R.S. Cotran, *The Lower Urinary Tract*, in *Robbins Pathologic Basis of Disease*, T.

Collins, Editor. 1999, W. B. Saunders Company: Philadelphia. p. 1003-1008). Currently, no reliable methods exist to screen and detect bladder cancer. Therefore, better methods to detect bladder tumors at early stage are necessary to decrease morbidity and mortality.

[0465] Over 95% of bladder tumors are epithelial in origin, with the remainder being

mesenchymal tumors. Approximately 90% of epithelial tumors are composed of transitional cells and are called transitional cell carcinoma (TCC), while the remaining 10% are squamous and glandular carcinomas. TCC is the fifth most common malignancy in the U.S. and second most common genitourinary carcinoma. TCC is segregated into two major categories: low-grade and high-grade. Low-grade TCCs are always papillary, noninvasive lesions that recapitulate normal transitional epithelium. These cases have an excellent prognosis. High-grade TCCs may be papillary, nodular, or both and exhibit considerable cellular pleomorphism and anaplasia. They account for 50% of bladder tumors, have metastatic potential, and are lethal in 60% of cases within 10 years of the diagnosis (Crawford, J.M. and R.S. Cotran, *The Lower Urinary Tract*, in *Robbins Pathologic Basis of Disease*, T. Collins, Editor. 1999, W. B. Saunders Company: Philadelphia. p. 1003-1008).

[0466] Urinary cytology is a conventional screening method and provides useful diagnostic information for high-grade bladder tumors (Koss, L.G., et al., Diagnostic value of cytology of voided urine. Acta Cytol, 1985. 29(5): p. 810-6). However, because of the lack of morphologic alterations in low-grade tumors, its efficacy in detection of low-grade papillary transitional cell carcinoma (TCC) is less reliable (Busch, C., et al., Malignancy grading of epithelial bladder tumours. Reproducibility of grading and comparison between forceps biopsy, aspiration biopsy and exfoliative cytology. Scand J Urol Nephrol, 1977. 11(2): p. 143-8; Murphy, W.M., et al., Urinary cytology and bladder cancer. The cellular features of transitional cell neoplasms. Cancer, 1984. 53(7): p. 1555-65; Shenoy, U.A., T.V. Colby, and G.B. Schumann, Reliability of urinary cytodiagnosis in urothelial neoplasms. Cancer, 1985. 56(8): p. 2041-5). Cystoscopy is invasive and bothersome to the patient. Exophytic tumors are reliably diagnosed by cystoscopy but flat TCC, particularly carcinoma in situ, remains an endoscopic dilemma (Lin, S., et al., Cytokeratin 20 as an immunocytochemical marker for detection of urothelial carcinoma in atypical cytology: preliminary retrospective study on archived urine slides. Cancer Detect Prev, 2001. 25(2): p. 202-9; Heicappell, R., et al., Evaluation of urinary bladder cancer antigen as a marker for diagnosis of transitional cell carcinoma of the urinary bladder. Scand J Clin Lab

Invest, 2000. 60(4): p. 275-82). Typically, both urinary cytology and cystoscopy are used together with biopsy, when necessary, to optimize diagnostic sensitivity. These tests, alone or combined, are insufficient for early detection or assessment of recurrence or disease progression. [0467] Immunocytochemistry (ICC) is gaining in popularity because it is noninvasive and has increased sensitivity compared to urine cytology. Urine tests for nuclear matrix proteins (NMP22) and human complement factor H-related proteins (BTA stat) have been on the market for several years, although they have had limited impact on the use of cystoscopy. NMP22 is a more sensitive test than BTA stat, but they both suffer from insufficient specificity and a falsepositive rate that is problematic (Ramakumar, S., et al., Comparison of screening methods in the detection of bladder cancer. J Urol, 1999. 161(2): p. 388-94; Ross, J.S. and M.B. Cohen. Detecting recurrent bladder cancer: new methods and biomarkers. Expert Rev Mol Diagn, 2001. 1(1): p. 39-52). However, FDA recently cleared an improved NMP22 assay, ImmunoCyt, from Matritech. This new assay contains two antibodies and works like the home pregnancy test. Its claims include a lower false-positive rate because the assay is not affected by the presence of blood in urine, and increased diagnostic accuracy in conjunction with cystoscopy. ImmunoCyt is a cocktail of three tumor markers labeled with fluorescent markers. It recognizes a mucin glycoprotein and a form of carcinoembronyonic antigen (CEA)-expressed by tumor cells in the bladder. However, it is used for monitoring, not screening, of recurrent bladder cancer and needs fluorescence microscopy for viewing (Mian, C., et al., Immunocyt: a new tool for detecting transitional cell cancer of the urinary tract. J Urol, 1999. 161(5): p. 1486-9).

[0468] Immunohistochemistry (IHC) is the most widely used evaluation method of bladder cancer for clinical urologists. Most tumor markers that have been studied and merit a role in the clinical decision-making process for bladder cancer have evolved from the application of IHC (Williams, S.G., M. Buscarini, and J.P. Stein, *Molecular markers for diagnosis, staging, and prognosis of bladder cancer*. Oncology (Huntingt), 2001. 15(11): p. 1461-70, 1473-4, 1476; discussion 1476-84). Unfortunately, none of the tumor markers for detecting bladder cancer is a "magic bullet" with both high sensitivity and specificity. Alternative ways to enhance diagnostic accuracy are necessary.

[0469] An alternative way to enhance diagnostic accuracy is to develop a panel comprising a plurality of probes each of which specifically binds a marker associated with bladder cancer.

Each candidate probe is to be tested by IHC or ICC. For ICC, the specimen will often be a urine

sample. The tradeoff of doing ICC, rather than IHC, is that urine cytological specimens usually have fewer cells than tissue sections. In some embodiments, high quality monoclonal or polyclonal antibodies may be used to assure assay good performance. In other embodiments, patients who are as close as possible in age, gender, diagnosis, tumor grade, tumor size, clinical stage, etc. may be studied. In further embodiments, the same patient may have urine collections as often as medically indicated and possible. Additionally, slides can be made by spiking tumor cells into cell suspension and testing before actual patient specimens are tested.

[0470] The specimen, either formalin-fixed paraffin-embedded (FFPE) tissue for IHC or urine cytology for ICC, will be obtained from medical institutions. Once the specimens are collected, the specimens will be processed and analyzed. Statistical analysis will be used to design panels, as described above for lung cancer. During processing, technical issues such as cell smears or pellets not sticking to slides during harsh washings may occur in some embodiments. However, such issues can readily be addressed by manipulation of software or modifying staining protocols to mitigate such problems. In some embodiments, the specimens will be processed and analyzed using a device that automatically samples the specimen and prepares monolayer slides for cyto-interpretation or diagnosis.

[0471] It is anticipated that a broad menu of probes will be used initially. The number of probes will be pruned to a suitably sized panel in order to retain a high level of sensitivity and specificity. Selection of the final probes will be based on a pre-defined threshold of the percentage of positive stained tumor cells. Sophisticated statistical analysis will be employed to make these determinations. Since the panel-assay approach to detecting malignancies is applicable to solid tumors, and several of the same tumor markers are in different panels, this method may be carried out in parallel, as well as serially. In this manner, the assay development process can be expedited.

[0472] The initial probes will be pruned to a suitably sized panel with high sensitivity and specificity. The selection of final probes is based on a pre-defined threshold of a percentage of positive stained tumor cells. Once a final panel of tests using IHC is determined, specimens will be tested by ICC. The established ICC probes will be tested on urine specimens as a panel. In some embodiments, automated staining will be employed, therefore, standardization can be achieved and results interpretation will be more consistent.

[0473] Compared with lung cancers, which have five subtypes (adenocarcinoma, squamous cell carcinoma, small cell carcinoma, large cell carcinoma, mesothelioma), over 90% of bladder cancer is categorized as transitional cell carcinoma (TCC). This allows the bladder tumor panel to be more specific at targeting one type of cancer. In addition, detection and diagnosis of bladder tumor can be FFPE-based and/or cell-based (urinary cytology).

Library of Probes/Markers

[0474] Various published scientific sources containing information about cancer markers were reviewed. An arbitrary criterion of 20% or greater positivity of bladder cancer was used to select probes for a preferred panel for detection and/or diagnosis of bladder cancer. The term "20% or greater positivity" means that if 100 tumor cases were studied, 20 or more of these cases would have shown a presence of the individual marker, while the remaining 80 or fewer cases would not have shown a presence of the individual marker. A preferred panel may include markers selected from BL2-10D1, C-erbB-2, CD44s Standard, Splice Variant CD44v6, Splice Variant CD44v3, Caveolin-1, Collagenase, Cyclin D1, Cyclooxygenase-1 (COX-1), Cyclooxygenase-2 (COX-2), Cytokeratin 20 (CK20), E-cadherin, Epidermal Growth Factor Receptor (EGFR), Heat Shock Protein-90 (HSP-90), IL-6, IL-10, HLA-DR, Human Mis-Match Repair Gene (hMSH2), Lewis X, MDM2, Nuclear Matrix Protein 22 (NMP-22), p53, PCNA, MIB1 (Ki-67), Retinoblastoma (Rb), Survivin, Transforming Growth Factor-β1, Transforming Growth Factor-β1 Receptor I, Transforming Growth Factor-β1 Receptor II and UBC (CK8 and CK18). A brief description of the library of probes/markers utilized in the present example is provided below.

BL2-10D1

[0475] This is an IgM antibody. A hypbridoma cell line secreting an IgM monoclonal antibody was produced after immunizing a mouse with RT4 cells and a suspension of human bladder carcinoma cells (Longin, A., et al., A monoclonal antibody (BL2-10D1) reacting with a bladder-cancer-associated antigen. Int J Cancer, 1989. 43(2): p. 183-9). It shows a strong reactivity with bladder tumors but not with normal urothelium except 5% to 10% of umbrella cells. This antibody reacts with most of the papillary Grade 1 and Grade 2 TCCs and with carcinoma in situ, whereas papillary Grade 3 and invasive non-papillary TCC show poor reactivity. Longin (Longin, A., et al., A useful monoclonal antibody (BL2-10D1) to identify tumor cells in urine cytology. Cancer, 1990. 65(6): p. 1412-7) and associates demonstrated that all urine from patients

with a low-grade (G1) tumors was stained with BL2-10D1. Grades 2 or 3 were not always stained. Such results suggest that BL2-10D1 may be considered a valuable marker of early detection of bladder cancer.

C-erbB-2

[0476] The c-erbB-2 gene encodes a transmembrane tyrosine kinase that is the receptor for a family of peptide hormones. C-erbB-2 amplification has been found in transitional cell carcinomas. Previous studies have observed an association of c-erbB-2 with metastasis, as well as with tumor grade or stage (Ioachim, E., et al., Immunohistochemical expression of retinoblastoma gene product (Rb), p53 protein, MDM2, c-erbB-2, HLA-DR and proliferation indices in human urinary bladder carcinoma. Histol Histopathol, 2000. 15(3): p. 721-7).

CD44 Standard (CD44s) and Splice Variants CD44v6 and CD44v3

[0477] CD44 is a transmembrane cell surface receptor. It has been associated with diverse functions, including cell-to-cell adhesion, cell matrix interaction, and tumor metastasis. The significance of CD44 isoforms in tumor development and its progression has been reported in various tumors. In a study by Masuda et al (Masuda, M., et al., Expression and prognostic value of CD44 isoforms in transitional cell carcinoma of renal pelvis and ureter. J Urol, 1999. 161(3): p. 805-8; discussion 808-9), expression of CD44s, CD44v6 and CD44v3 was significantly decreased in relation to histologic grade of bladder cancer. However, all of these isoforms were expressed strongly on the cytoplasmic membrane of basal cells of normal urothelial mucosa. However, the superficial layers of normal urothelial mucosa did not express them.

Caveolin-1

[0478] Caveolae are abundant in numerous cell types, ranging from adipocytes and endothelial cells to type I pneumocytes and skeletal muscle cells. Three constituent caveolin protein family members have been identified, caveolin-1, caveolin-2 and caveolin-3. CAV-1 gene has been mapped to chromosome 7q31 and has much scientific interest as a potential site of tumor suppressor activity. It presumably involves signal transduction by interacting with a broad range of signal transducing molecules and receptors (Src, G protein, and EGFR). Rajjayabun (Ioachim, E., et al., Immunohistochemical expression of retinoblastoma gene product (Rb), p53 protein, MDM2, c-erbB-2, HLA-DR and proliferation indices in human urinary bladder carcinoma. Histol Histopathol, 2000. 15(3): p. 721-7) first showed CAV-1 immunoreactivity in high-grade bladder cancers.

Collagenase

[0479] Collagenase is known to dissolve collagen and repair and maintain tissues. It is secreted from epithelial cells, neutrophils, histiocytes and fibroblasts. Increased expression of collagenase has been associated with breast and thyroid cancer. RT-PCR study showed increased mRNA in urothelial carcinoma and one study demonstrated 34% and 45% of patients with TCC showed positive expression of collagenase on cytologic and histologic specimens, respectively. No expression was found on benign lesions (Hattori, M., E. Ohno, and H. Kuramoto, Immunocytochemistry of collagenase expression in transitional cell carcinoma of the bladder. Acta Cytol, 2000. 44(5): p. 771-7).

Cyclin D1

[0480] Cyclin D1 gene product contributes to the regulation of the G1/S-phase transition of cell cycle and is a candidate oncogen. It has been shown to correlate with low-grade, low-stage and papillary tumor growth in primary bladder carcinomas and it has been suggested to play an important role in bladder cancer progression (Byrne, R.R., et al., E-cadherin immunostaining of bladder transitional cell carcinoma, carcinoma in situ and lymph node metastases with long-term followup. J Urol, 2001. 165(5): p. 1473-9).

Cyclooxygenase-1 (Cox-1) and Cyclooxygenase-2 (Cox-2)

[0481] Cyclooxygenases are the rate-limiting enzymes catalyzing the initial step in the formation of prostaglandins that are involved in inflammation, immune responses, mitogenesis and apoptosis. Cyclooxygenase-1 (Cox-1) is constitutively expressed in most tissues at a rather stable level. The low basal activity of the inducible form, cyclooxygenase-2 (Cox-2), is increased during inflammatory processes by cytokines, growth factors, oncogenes and tumor promoters. Increased cyclooxygenase activity and, consequently, elevated prostaglandin levels have been observed in gastroenterological malignancies, as well as bladder cancer. Bostrom et al (Bostrom, P.J., et al., Expression of cyclooxygenase-1 and -2 in urinary bladder carcinomas in vivo and in vitro and prostaglandin E2 synthesis in cultured bladder cancer cells. Pathology, 2001. 33(4): p. 469-74) showed diffuse, moderate to strong, cytoplasmic immunosignal for Cox-2 that was detected in all 29 TCCs of bladder. Normal urothelium in the specimen also stained for Cox-2, but the intensity was weak. Cox-1 staining was detected in tissue from 18 of 29 TCC specimens (62%) but the signal was weak in 16 of the 18 specimens. Immunosignal from Cox-1 was detected only in a few normal specimens and was weak to moderate.

Cytokeratin 20 (CK 20)

[0482] Cytokeratin 20 (CK 20), one of 20 known cytokeratins, is a constituent of the intermediate filaments of epithelial cells. IHC study has shown CK 20 was expressed in urothelial cells of patients with urothelial carcinoma or urothelial dysplasia. In normal urothelium, CK 20 expression was restricted on superficial umbrella cells. By using ICC, Lin (Lin, S., et al., Cytokeratin 20 as an immunocytochemical marker for detection of urothelial carcinoma in atypical cytology: preliminary retrospective study on archived urine slides. Cancer Detect Prev, 2001. 25(2): p. 202-9) and associates showed CK 20 was positive in 95% of bladder cancer patients and only positive in 10% of normal control.

E-cadherin

[0483] E-cadherin is expressed in all epithelial tissue and is found on the plasma membrane of squamous and transitional cells. E-cadherin mediated cell adhesion is involved in tumor progression and metastasis. IHC studies of E-cadherin in transitional cell carcinoma of the bladder have demonstrated a significant association of aberrant E-cadherin expression with advanced tumor stage and loss of differentiation. Byrne et al (Byrne, R.R., et al., E-cadherin immunostaining of bladder transitional cell carcinoma, carcinoma in situ and lymph node metastases with long-term followup. J Urol, 2001. 165(5): p. 1473-9) showed 59 (77%) bladder tumors had loss of normal membrane E-cadherin, whereas preserved E-cadherin expression was seen in normal urothelium.

Epidermal Growth Factor Receptor (EGFR)

[0484] Epidermal growth factor is a potent mitogen and its actions are mediated by binding to the external domain of epidermal growth factor receptor (EFGR). EGFR is a transmembrane protein receptor with tyrosine kinase activity. The cytoplasmic and internal domains of EGFR have close similarity with the oncogene product of the avian erythroblastosisi virus (v-erb-B-2). Increased levels of EGFR are found in solid tumors, including squamous cell carcinoma of lung, head, neck, cervix, breast, prostate and bladder (Neal, D.E., et al., *The epidermal growth factor receptor and the prognosis of bladder cancer*. Cancer, 1990. 65(7): p. 1619-25). The range of positivity of bladder cancer is 31-48% (American Cancer Society. 2000).

Heat Shock Protein-90 (HSP-90), IL-6 and IL-10

[0485] During the early stages of bladder cancer, a cascade of immunological reactions takes place and various proteins, including heat shock protein (HSP-90) and cytokines, are secreted in

large amounts. HSP-90 is one of the most important members of the HSP family. Of the 56 bladder carcinoma studied by IHC, 52 (93%) expressed HSP-90, 48 (86%) expressed IL-6 and 45 (80%) expressed IL-10. High-grade and muscle-invasive tumors contained significantly higher levels of HSP-90 and IL-6 than low-grade tumors. Normal urothelium adjacent to tumor areas do not show HSP-90 staining. IL-6 and IL-10 showed scarce immunoreactivity (Cardillo, M.R., P. Sale, and F. Di Silverio, *Heat shock protein-90, IL-6 and IL-10 in bladder cancer*. Anticancer Res, 2000. 20(6B): p. 4579-83). This suggests IL-6 and IL-10 may be turned on at a relatively low stage during tumor development.

HLA-DR

[0486] Although the functional role of HLA-DR is to mediate communication among immuno-competent cells, it also has been shown that HLA-DR antigen expression is independent of lymphocyte subpopulations in bladder cancer (Ioachim, E., et al., *Immunohistochemical expression of retinoblastoma gene product (Rb)*, p53 protein, MDM2, c-erbB-2, HLA-DR and proliferation indices in human urinary bladder carcinoma. Histol Histopathol, 2000. 15(3): p. 721-7).

Human Mis-Match Repair Gene (hMSH2)

[0487] This is the prototype mismatch repair gene (MMR). hMHS2 recognizes and binds to nucleotide mismatches and in conjunction with other MMR proteins directs the coordinated correction of DNA replication errors. Mutation of MMR genes has been reported in 40% of hereditary non-polyposis colon cancer (HNPCC) and recent IHC studies suggest that bladder tumors with decreased expression of hMSH2 are associated with higher grade and recurrence. One study showed all bladder tumors are positive for hMSH2, whereas normal bladder mucosa are negative by IHC (Leach, F.S., et al., Expression of the human mismatch repair gene hMSH2: a potential marker for urothelial malignancy. Cancer, 2000. 88(10): p. 2333-41).

Lewis X

[0488] ABH and Lewis blood group-related antigens are present on the surface of normal urothelium. The Lewis X antigen is normally absent from urothelial cells in the adult, except for occasional umbrella cells. Sheinfeld (Sheinfeld, J., et al., Enhanced bladder cancer detection with the Lewis X antigen as a marker of neoplastic transformation. J Urol, 1990. 143(2): p. 285-8) and associates have used immunostaining of the Lewis X antigen on epithelial cells from bladder washing specimens for detection of bladder tumors and reported sensitivity of 86% and

specificity of 87%. Golijanin (Golijanin, D., et al., Detection of bladder tumors by immunostaining of the Lewis X antigen in cells from voided urine. Urology, 1995. 46(2): p. 173-7) and associates also showed high sensitivity and specificity of Lewis X immunostaining of urine samples. High-grade and low-grade transitional cell tumors were detected with equal efficiency.

MDM2

[0489] This is a cellular proto-oncogene product. MDM2 has been shown to bind to p53 and acts as a negative regulator, inhibiting its transcriptional trans-activation. It has been shown that aberrant MDM2 and p53 phenotypes may be important diagnostic markers in bladder cancer patients (Ioachim, E., et al., Immunohistochemical expression of retinoblastoma gene product (Rb), p53 protein, MDM2, c-erbB-2, HLA-DR and proliferation indices in human urinary bladder carcinoma. Histol Histopathol, 2000. 15(3): p. 721-7).

Nuclear Matrix Protein 22 (NMP-22)

[0490] This nuclear matrix protein plays an important role in the structural framework of the nucleus, in DNA replication and in gene expression. Significantly increased concentrations of NMPs have been found with neoplastic transformation and in carcinomas of the breast, colon and bladder. Soluble NMPs can be detected in the urine from bladder cancers using antibodies against select epitopes of NMP (NMP-22). Landman et al (Landman, J., et al., Sensitivity and specificity of NMP-22, telomerase, and BTA in the detection of human bladder cancer. Urology, 1998. 52(3): p. 398-402) found the overall sensitivity to be 81%.

p53

[0491] This human tumor suppressor gene encodes a nuclear phosphoprotein that facilitates DNA repair after genomic damage. Wild type p53 degrades. Mutant p53 does not and therefore accumulates in the cell. Mutant p53 can be detected by IHC. Several studies associate p53 mutation with high-grade bladder cancer and unfavorable prognosis (Vollmer, R.T., et al., Invasion of the bladder by transitional cell carcinoma: its relation to histologic grade and expression of p53, MIB-1, c-erb B-2, epidermal growth factor receptor, and bcl-2. Cancer, 1998. 82(4): p. 715-23; Nakopoulou, L., et al., The prevalence of bcl-2, p53, and Ki-67 immunoreactivity in transitional cell bladder carcinomas and their clinicopathologic correlates. Hum Pathol, 1998. 29(2): p. 146-54; Vorreuther, R., et al., Expression of immunohistochemical markers (PCNA, Ki-67, 486p and p53) on paraffin sections and their relation to the recurrence

rate of superficial bladder tumors. Urol Int, 1997. 59(2): p. 88-94; Li, B., et al., Reciprocal expression of bcl-2 and p53 oncoproteins in urothelial dysplasia and carcinoma of the urinary bladder. Urol Res, 1998. 26(4): p. 235-41). The reported p53 mutation rate in bladder cancer varies from 30% to 58%. Wu et al (Wu, T.T., et al., The role of bcl-2, p53, and ki-67 index in predicting tumor recurrence for low grade superficial transitional cell bladder carcinoma. J Urol, 2000. 163(3): p. 758-60) demonstrated 42 of 60 (70%) cases had p53 mutation.

PCNA and MIB1 (Ki-67)

[0492] Both are cell proliferation nuclear markers. They are expressed in a variety of tumors including bladder cancer. Studies demonstrated a strong correlation between PCNA and MIB1, suggesting either of the markers could be used (Ioachim, E., et al., Immunohistochemical expression of retinoblastoma gene product (Rb), p53 protein, MDM2, c-erbB-2, HLA-DR and proliferation indices in human urinary bladder carcinoma. Histol Histopathol, 2000. 15(3): p. 721-7; Lavezzi, A.M., L. Terni, and L. Matturri, PCNA immunostaining as a valid alternative to tritiated thymidine-autoradiography to detect proliferative cell fraction in transitional cell bladder carcinomas. In Vivo, 2000. 14(3): p. 447-51).

Retinoblastoma (Rb)

[0493] Rb is a tumor suppressor gene and may play a role in the initiation and progression of human tumors. Rb codes for nuclear phosphoproteins present in normal cells and are thought to be involved in cell cycle control and in negative regulation of cell growth. Alterations of the Rb gene have been observed in several epithelial tumors suggesting structural abnormalities, including mutations and/or deletions of this gene, may result in the inactivation of tumor suppressor protein and may be involved in tumorigenesis.

Survivin

[0494] Survivin is a 142 amino acid protein. It is expressed in the G2/M phase of the cell cycle and associates with microtubules of the mitotic spindle. It is an inhibitor of apoptosis that is selectively over-expressed in common human cancers, but not in normal tissues, and correlates with aggressive disease and unfavorable outcomes (Lehner, R., et al., Immunohistochemical localization of the IAP protein survivin in bladder mucosa and transitional cell carcinoma. Appl Immunohistochem Mol Morphol, 2002. 10(2): p. 134-8).

Transforming Growth Factor-\$1, and Receptors I and II

[0495] TGF-β1 is a pleiotropic growth factor regulating cellular proliferation, differentiation, migration, immune response, angiogenesis and apoptosis. It has been shown to inhibit normal epithelial cell growth by reducing the ability of cells to enter S-phase. Conversely, many carcinoma cells show resistance to growth inhibition by TGF-β1. The effects of TGF-β1 are mediated by membrane-bound serine-threonine kinase receptors, currently referred to as TGF-β1 receptor I (TGF-β1I) and TGF-β1 receptor II (TGF-β1II). The loss of expression of receptors I and II has been found in several cancers including prostate, colon, ovarian, and bladder. In one study, expression of TGF-β1, TGF-β1I, and TGF-β1II was altered in 51 (64%), 34 (43%), and 38 (48%) of bladder cancer cases. Over-expression of TGF-β1 was seen in high-grade tumors and is negative in normal bladder mucosa. Loss of TGF-β1I and β1II was associated with invasive tumor stage (Kim, J.H., et al., *Predictive value of expression of transforming growth factor-beta(1) and its receptors in transitional cell carcinoma of the urinary bladder.* Cancer, 2001. 92(6): p. 1475-83).

UBC (CK8 and CK18)

[0496] This is an enzyme immunoassay that measures the concentration of cytokeratin fragments, 8 and 18, in the urine. These cytokeratins are co-expressed by simple epithelia and carcinomas, among them normal urothelium and TCC. Heicappell et al demonstrated sensitivities of UBC to range from 22% and 75% with the overall specificities of 77% (Heicappell, R., et al., Evaluation of urinary bladder cancer antigen as a marker for diagnosis of transitional cell carcinoma of the urinary bladder. Scand J Clin Lab Invest, 2000. 60(4): p. 275-82).

Tissue-Based Markers

[0497] A tissue-based panel for detecting and/or diagnosing bladder cancer includes diagnostic and prognostic markers, as well as those associated with metastatic potential, tumor grade and stage determinations. It has potential value not only in patient screening and diagnosis, but also in patient stratification, to guide clinical therapy. Tissue-based panels can also be tested on urine specimens to see whether the results are superior to the existing cell-based panel. A preferred tissue-based panel for detecting and/or diagnosing bladder cancer comprises markers selected

from p53, Rb, MDM2, EGFR and survivin. The importance of these tumor markers with respect to bladder cancer is detailed below.

[0498] Detection of high-grade bladder cancer usually is not a problem because severe cytological atypia is present. On the other hand, detection of low-grade tumors is a continuing challenge for the cytopathologist. Understanding the pathogenesis of a bladder tumor is crucial to pinpointing the earliest events during tumorigenesis. Cigarette smoking, exposure to arylamines, long-term use of analgesics, and exposure to cyclophosphamide increase the risk of bladder cancer. How these influences induce cancer is unclear, but a number of genetic alterations have been observed in transitional cell carcinoma. The evidence suggests chromosome deletions involving tumor-suppressor genes and subsequent loss of cell-cycle control are important in the early development of bladder tumor (Crawford, J.M. and R.S. Cotran, *The Lower Urinary Tract*, in *Robbins Pathologic Basis of Disease*, T. Collins, Editor. 1999, W. B. Saunders Company: Philadelphia. p. 1003-1008; Williams, S.G., M. Buscarini, and J.P. Stein, *Molecular markers for diagnosis, staging, and prognosis of bladder cancer*. Oncology (Huntingt), 2001. 15(11): p. 1461-70, 1473-4, 1476; discussion 1476-84).

[0499] Between 30% to 60% of bladder tumors have chromosome 9 monosomy or deletions of 9p and 9q, as well as deletions of 17p, 13q, 11p and 14q (Gibas, Z. and L. Gibas, Cytogenetics of bladder cancer. Cancer Genet Cytogenet, 1997. 95(1): p. 108-15). Chromosome 9 deletions are the only genetic changes frequently present in superficial papillary tumors and occasionally in noninvasive flat tumors. The 9p deletion, 9p21, involves tumor-suppressor gene p16. However, studies demonstrate p16 protein expression is not decreased in 9p deletions. On the other hand, many invasive transitional cell carcinomas show deletions of 17p and mutations in the p53 gene, suggesting that alterations in p53 contribute to the progression of transitional cell carcinoma. [0500] Mutations in p53 are also found in flat in situ bladder cancer. MDM2, a protooncogene, plays a pivotal role in regulation of the cell cycle by binding with p53. MDM2 can also interact with other critical elements of the cell cycle and apoptotic regulatory controls, such as E2F and Rb (Martin, K., et al., Stimulation of E2F1/DP1 transcriptional activity by MDM2 oncoprotein. Nature, 1995. 375(6533): p. 691-4; Xiao, Z.X., et al., Interaction between the retinoblastoma protein and the oncoprotein MDM2. Nature, 1995. 375(6533): p. 694-8). The 13q deletion is that of the retinoblastoma gene (Rb) and mutation of the Rb gene is considered to be of central importance in the pathogenesis of many human malignant neoplasms. Recently, it

has become clear that the genomic loss of pRb leads to uncontrolled cell growth and apoptosis (i.e., cell death). Previous studies in bladder cancer have reported that inactivation of Rb is associated with tumor progression (Ioachim, E., et al., Immunohistochemical expression of retinoblastoma gene product (Rb), p53 protein, MDM2, c-erbB-2, HLA-DR and proliferation indices in human urinary bladder carcinoma. Histol Histopathol, 2000. 15(3): p. 721-7).

[0501] The recurrent rate is high for low-grade bladder cancer. For patients without muscle invasion, the long-term outlook is better but more than half have recurrent tumors develop within the bladder that are usually similar in stage and grade to the primary cancer. EGFR has been shown to be associated with increased recurrent rate and the progression of superficial to invasive bladder cancer (Neal, D.E., et al., The epidermal growth factor receptor and the prognosis of bladder cancer. Cancer, 1990. 65(7): p. 1619-25; Izawa, J.I., et al., Differential expression of progression-related genes in the evolution of superficial to invasive transitional cell carcinoma of the bladder. Oncol Rep, 2001. 8(1): p. 9-15).

[0502] Survivin is a new bladder tumor marker expressed in the G2/M phase of the cell cycle and associates with microtubules of the mitotic spindle (Li, B., et al., Reciprocal expression of bcl-2 and p53 oncoproteins in urothelial dysplasia and carcinoma of the urinary bladder. Urol Res, 1998. 26(4): p. 235-41). Disruption of survivin-microtubule interactions results in the loss of survivin's anti-apoptotic function and in activation of caspase-3. Caspase-3 is an essential death factor for the Fas-mediated cell death and its inactivation in cells is initiated by an interaction with p21 (Tamm, I., et al., IAP-family protein survivin inhibits caspase activity and apoptosis induced by Fas (CD95), Bax, caspases, and anticancer drugs. Cancer Res, 1998. 58(23): p. 5315-20). Survivin is also involved in cell-cycle control. More importantly, studies show that nuclear staining of survivin is present in a majority of transitional cell carcinoma and none in healthy bladder mucosa (Lehner, R., et al., Immunohistochemical localization of the IAP protein survivin in bladder mucosa and transitional cell carcinoma. Appl Immunohistochem Mol Morphol, 2002. 10(2): p. 134-8).

Cell-Based Immunocytochemical Markers

[0503] The cell-based panel has markers specific for early stage, low-grade TCC, as well as for late, high-grade TCC. It can be used to screen and detect early stage, low-grade TCC. For difficult cases, a cytoscopic biopsy can be followed to confirm the diagnosis with the tissue-based panel. It can also be used to diagnose high-grade TCC if a patient presents at late stage.

The results can guide therapeutic decisions by urologists. A preferred cell-based panel for detecting and/or diagnosing bladder cancer using immunocytochemical techniques comprises markers selected from BL2-10D1, cytokeratin 20, Lewis X and NMP-22. The importance of these tumor markers with respect to bladder cancer is detailed below.

[0504] The introduction of novel molecular markers into clinical urology has increased the diagnostic accuracy of bladder cancer compared to conventional urinary cytology. A number of commercial products, as well as individual markers, are available. Each marker, however, exhibits a different sensitivity and specificity. Furthermore, no single immunocytochemical assay has yet replaced invasive cystoscopy.

[0505] Uniqueness is evident with several markers. For example, BL2-10D1 has a consistent reactivity with low-grade TCC versus its non-consistent reactivity with high-grade TCC (Longin, A., et al., A useful monoclonal antibody (BL2-10D1) to identify tumor cells in urine cytology. Cancer, 1990. 65(6): p. 1412-7). CK 20 has an overall sensitivity of 94% and specificity of 80% for the detection of TCC (Lin, C.W., et al., Detection of tumor cells in bladder washings by a monoclonal antibody to human bladder tumor-associated antigen. J Urol, 1988. 140(3): p. 672-7), and in benign epithelium staining is limited on the surface of umbrella cells while in dysplastic epithelium the staining involves full thickness (Harnden, P., et al., Cytokeratin 20 as an objective marker of urothelial dysplasia. Br J Urol, 1996. 78(6): p. 870-5). Immunostaining of the Lewis X antigen on epithelial cells from bladder washing specimens found a sensitivity of 86% and specificity of 87% (Sheinfeld, J., et al., Enhanced bladder cancer detection with the Lewis X antigen as a marker of neoplastic transformation. J Urol, 1990. 143(2): p. 285-8). And, studies done to compare NMP-22, telomerase, and BTA assay demonstrate that NM-22 has the highest sensitivity and specificity (Landman, J., et al., Sensitivity and specificity of NMP-22, telomerase, and BTA in the detection of human bladder cancer. Urology, 1998. 52(3): p. 398-402; Friedrich, M.G., et al., Clinical use of urinary markers for the detection and prognosis of bladder carcinoma: a comparison of immunocytology with monoclonal antibodies against Lewis X and 486p3/12 with the BTA STAT and NMP22 tests. J Urol, 2002. 168(2): p. 470-4). [0506] By combining biomarkers for one or more panel-assays, sensitivities and specificities increase significantly (Eissa, S., et al., Comparative evaluation of the nuclear matrix protein, fibronectin, urinary bladder cancer antigen and voided urine cytology in the detection of bladder tumors. J Urol, 2002. 168(2): p. 465-9). An ideal bladder cancer assay should be non-invasive,

sensitive, specific, and cost effective. While the tissue-based panel and cell-based panels may be used alone, markers from a tissue-based panel and from a cell-based panel can be combined again to have a final optimized panel.

[0507] The following abbreviations were used throughout this example: CD44s = CD44
Standard, CD44v6 = CD44 splice variant 6, CD44v3 = CD44 splice variant 3, CK = Cytokeratin,
Cox-1 = Cyclooxygenase-1, Cox-2 = Cyclooxygenase-2, EGFR = Epidermal Growth Factor
Receptor, ELISA = Enzyme Linked ImmunoSorbent Assay, FFPE = Formalin Fixed Paraffin
Embedded, FNA = Fine Needle Aspiration, FNB = Fine Needle Biopsy, HLA-DR = Human
Leukocyte Antigen-DR, hMSH2 = Human Mismatch Repair Gene, HNPCC = Hereditary
Nonpolyposis Colon Cancer, HSP-90 = Heat Shock Protein-90, IL-6 = Interlukin-6, IL-10 =
Interlukin-10, ICC = Immunocytochemistry, IHC = Immunohistochemistry, ISH = In Situ
Hybridization, NMP22 = Nuclear matrix proteins, PCNA = Proliferating Cell Nuclear Antigen,
PCR = Polymerase Chain Reaction, TCC = Transitional Cell Carcinoma, TGF = Transforming
Growth Factor, STDs = Sexual Transmitted Diseases and Rb = Retinoblastoma.

IV. Prostate Cancer

[0508] A variety of prostate tumor markers have been discovered to aid physicians in making timely, precise diagnoses, and to provide significantly better patient management. Unfortunately, none of these prostate tumor markers is a "magic bullet" with both high sensitivity and specificity. Therefore, alternative ways to enhance diagnostic accuracy are necessary. [0509] An alternative way to enhance diagnostic accuracy is to develop a panel comprising a plurality of probes each of which specifically binds a marker associated with prostate cancer. All candidate probes are to be tested with ICC and/or IHC techniques. In some embodiments, specimens may be obtained from a fine needle biopsy. Fine needle aspiration (FNA) is usually used for superficial and external organs, whereby a cell-suspension cytology specimen can be obtained, such as from breast, thyroid or a similar soft tissue mass. In the case of deep and internal organs, such as prostate, kidney or liver, a fine needle biopsy (FNB) technique is often used which may be guided by ultrasound or CT, and a tissue specimen is obtained. With the prostate gland, ultrasound is often used and a FNB is performed transrectally. In other embodiments, touch preparations (touch preps) of FNBs can be performed to generate an imprint of cells on a glass slide from tissue. In the case of a fluid or a bloody specimen from a FNB procedure, the cell suspension specimen can be generated.

[0510] Once the specimens are collected, the specimens will be processed and analyzed. Statistical analysis will be used to design panels, as described above for lung cancer. During processing, technical issues such as cell smears or pellets not sticking to slides during harsh washings may occur in some embodiments. However, such issues can readily be addressed by manipulation of software or modifying staining protocols can mitigate such problems. In some embodiments, the specimens will be processed and analyzed using a device that automatically samples the specimen and prepares slides for diagnosis. It is anticipated that a broad menu of probes will be used initially. The number of probes will be pruned to a suitably sized panel in order to retain a high level of sensitivity and specificity. Selection of the final probes will be based on a pre-defined threshold of the percentage of positive stained tumor cells. Sophisticated statistical analysis will be employed to make these determinations. Since the panel-assay approach to detecting malignancies is applicable to solid tumors, and several of the same tumor markers are in different panels, this method may be carried out in parallel, as well as serially. In this manner, the assay development process can be expedited.

Library of Probes/Markers

[0511] Various sources containing information about cancer markers were reviewed. An arbitrary criterion of 20% or greater positivity of prostate cancer was used to select probes for a preferred panel for detection and/or diagnosis of prostate cancer. The term "20% or greater positivity" means that if 100 tumor cases were studied, 20 or more of these cases would have shown a presence of the individual marker, while the remaining 80 cases would not have shown a presence of the individual marker. A preferred panel may include molecular markers selected from 34βE12, B72.3, C-erb.B2, E-Cadherin, Epidermal Growth Factor Receptor (EGFR), Fatty Acid Senthase (FAS), ID-1, Kallikrein 2, Ki-67, Leu 7, MDM2, N-Cadherin, P504S, p53, Prostate Acid Phosphatase (PAP), Prostate Inhibin Peptide (PI) and Prostate Specific Antigen (PSA). A brief description of the library of probes/markers utilized in the present example is provided below.

34BE12

[0512] This is a high-molecular-weight cytokeratin (HMWCK) that is present in basal cells of the prostate. Adenocarcinoma of the prostate lacks imunoreactivity with this antibody. Yang and associates (Yang, X.J., et al., Rare expression of high-molecular-weight cytokeratin in adenocarcinoma of the prostate gland: a study of 100 cases of metastatic and locally advanced

prostate cancer. Am J Surg Pathol, 1999. 23(2): p. 147-52) studied 100 cases of metastatic and locally advanced prostate cancer with 34βE12 by immunohistochemistry. Only 4 cases were positive for 34βE12. They concluded 34βE12 is a very useful marker to differentiate prostate adenocarcinoma from normal prostatic glands. Another group studied 228 equivocal cases and found 34βE12 of great value in confirming, establishing, or changing diagnoses in questionable foci seen in the everyday practice of surgical pathology (Wojno, K.J. and J.I. Epstein, The utility of basal cell-specific anti-cytokeratin antibody (34 beta E12) in the diagnosis of prostate cancer. A review of 228 cases. Am J Surg Pathol, 1995. 19(3): p. 251-60).

B72.3

[0513] This is a monoclonal antibody that recognizes tumors associated with glycoprotein 72. It has been shown to react with various types of adenocarcinoma, including prostate adenocarcinoma. Mazur and associates (Mazur, M.T. and J.J. Shultz, *Prostatic adenocarcinoma*. Evaluation of immunoreactivity to monoclonal antibody B72.3. Am J Clin Pathol, 1990. 93(4): p. 466-70) showed that over 95% of benign prostate disease was negative for B72.3, whereas 77% of prostate adenocarcinoma was at least focally positive and 100% well-differentiated prostate adenocarcinoma was B72.3 positive. Another study demonstrated that benign epithelium and stromal tissue did not immuno-stain with B72.3. Immunostaining was detected within the malignant cells in 30% of localized prostate adenocarcinomas (Myers, R.B., et al., Expression of tumor-associated glycoprotein 72 in prostatic intraepithelial neoplasia and prostatic adenocarcinoma. Mod Pathol, 1995. 8(3): p. 260-5).

C-erb-B2

[0514] This is an oncogene also known as Her2/neu, and its product is called oncoprotein. C-erb-B2 belongs to the epidermal growth factor receptor (EGFR) family that includes c-erb-B1, c-erb-B2, and c-erb-B3. Over-expression of c-erb-B2 is well known in breast carcinoma. The role of c-erb-B2 remains uncertain in the pathogenesis and progression of human prostate cancer. Previous studies have reported widely divergent rates of c-erb-B2 expression in primary prostate tumors, probably due to significant methodologic differences in the studies. Reese and associates (Reese, D.M., et al., HER2 protein expression and gene amplification in androgen-independent prostate cancer. Am J Clin Pathol, 2001. 116(2): p. 234-9) studied Her2 protein expression on androgen-independent prostate cancer by immunohistochemical and fluorescence in situ hybridization (FISH). They discovered that 36% of the specimens were positive for Her2 but

only 6% had gene amplification, indicating the mechanism for Her2 amplification and protein expression is different than in breast cancer. Another study demonstrated c-erb-B2 is a very useful biomarker for advanced prostate carcinoma (Arai, Y., T. Yoshiki, and O. Yoshida, c-erbB-2 oncoprotein: a potential biomarker of advanced prostate cancer. Prostate, 1997. 30(3): p. 195-201).

E-Cadherin

[0515] This is a transmembrane Ca++ dependent cell adhesion molecule. Decreased expression of E-Cadherin has been demonstrated in a number of carcinomas and has been associated with both a higher tendency to metastasize and a decreased survival rate. Kuniyasu and associates (Kuniyasu, H., et al., Relative expression of type IV collagenase, E-cadherin, and vascular endothelial growth factor/vascular permeability factor in prostatectomy specimens distinguishes organ-confined from pathologically advanced prostate cancers. Clin Cancer Res, 2000. 6(6): p. 2295-308) showed that a decreased expression of E-Cadherin was associated with an increasing Gleason score. Other studies also revealed E-Cadherin is down-regulated in prostatic bone metastases and reduced E-Cadherin is associated with poor prognosis in patients with prostate cancer (Bryden, A.A., et al., E-cadherin and beta-catenin are down-regulated in prostatic bone metastases. BJU Int, 2002. 89(4): p. 400-3; Umbas, R., et al., Decreased E-cadherin expression is associated with poor prognosis in patients with prostate cancer. Cancer Res, 1994. 54(14): p. 3929-33).

EGFR

[0516] Epidermal growth factor receptor (EGFR) is a transmembrane glycoprotein. It plays an important role in cell growth and differentiation. Epidermal growth factor (EGF), one of the ligands, interacts with cell-surface epidermal growth factor receptors (EGFR) to induce receptor tyrosine phosphorylation and activation of the intracellular signal-transduction pathways. EGF appears to be the predominant EGF-related growth factor in the normal prostate and in benign prostatic hyperplasia (BPH). EGFR is located in the basal/neuroendocrine (NE) compartment of the benign prostate and exhibits relatively androgen-independent expression. EGF-related peptides and EGFR are also present in neoplastic prostatic tissues. Androgen-independent cancer cells exhibit more EGFR expression and phos-phorylation than do androgen-responsive prostate cancer cells (Sherwood, E.R. and C. Lee, Epidermal growth factor-related peptides and the epidermal growth factor receptor in normal and malignant prostate. World J Urol, 1995. 13(5):

p. 290-6). One study found 22 of 46 prostate adenocarcinomas expressed EGFR (Cohen, D.W., et al., Expression of transforming growth factor-alpha and the epidermal growth factor receptor in human prostate tissues. J Urol, 1994. 152(6 Pt 1): p. 2120-4).

Fatty Acid Synthase (FAS)

[0517] Fatty acid synthase (FAS), or onco-antigen 519 (OA-519), is a key lipogenic enzyme. It has been recently associated with poor prognosis in breast cancers. Shurbaji and associates (Shurbaji, M.S., J.H. Kalbfleisch, and T.S. Thurmond, Immunohistochemical detection of a fatty acid synthase (OA-519) as a predictor of progression of prostate cancer. Hum Pathol, 1996. 27(9): p. 917-21) demonstrated that expression of FAS was in 57% of prostate cancers and FAS positive cancers were more likely to progress than FAS negative cancers. In Swinnen's study (Swinnen, J.V., et al., Overexpression of fatty acid synthase is an early and common event in the development of prostate cancer. Int J Cancer, 2002. 98(1): p. 19-22), benign hyperplastic glandular structures were all negative for FAS staining, immunohistochemical signal was evident in 24 of 25 low grade prostatic epithelial neoplasia (PIN) lesions, in 26 of 26 high grade PIN lesions and in 82 of 87 invasive carcinomas. Staining intensity tended to increase from low grade to high grade PIN to invasive carcinoma. Cancers with a high FAS expression had an overall high proliferative index. Another study found that FAS is a significant prognostic marker for prostate cancer and is one of the few markers that provides additional predictive information beyond that of the Gleason score (Epstein, J.I., M. Carmichael, and A.W. Partin, OA-519 (fatty acid synthase) as an independent predictor of pathologic state in adenocarcinoma of the prostate. Urology, 1995. 45(1): p. 81-6).

ID-1

[0518] The helix-loop-helix protein ID-1 serves to prevent basic helix-loop-helix transcription factors from binding to DNA, thus, inhibiting the transcription of differentiation associated genes. Over expression of ID-1 has been reported in certain tumors, such as breast, esophageal, pancreatic and medullary thyroid cancers. Ouyang and associates (Ouyang, X.S., et al., Over expression of ID-1 in prostate cancer. J Urol, 2002. 167(6): p. 2598-602) documented that negative to weak expression of ID-1 in normal prostate or BPH tissue was observed on immunohistochemical study and in situ hybridization. In contrast, all prostate cancer biopsies showed significant positive ID-1 expression in tumor cells at the messenger RNA and protein levels. Furthermore, expression of ID-1 was stronger in poorly differentiated than in well-

differentiated carcinomas, suggesting that the level of ID-1 expression may be associated with tumor grade.

Kallikrein 2

[0519] Human glandular kallikrein 2 (hK2) is a member of a multigene family of serine proteases. It is expressed in prostate epithelium and is under androgen regulation. Kallikrein 2 is present in serum and seminal fluid. It can form complexes with endogenous protease inhibitors (e.g., alpha2-macroglobulin and alpha1-antichymo-trypsin). Studies show the specificity of prostate cancer detection increased when kallikrein 2 combined with prostate-specific antigen (PSA) (Partin, A.W., et al., Use of human glandular kallikrein 2 for the detection of prostate cancer: preliminary analysis. Urology, 1999. 54(5): p. 839-45). Another study showed that the combination of human glandular kallikrein and free prostate-specific antigen (PSA) enhances discrimination between prostate cancer and benign prostatic hyperplasia in patients with a moderately increased total PSA (Magklara, A., et al., The combination of human glandular kallikrein and free prostate-specific antigen (PSA) enhances discrimination between prostate cancer and benign prostatic hyperplasia in patients with moderately increased total PSA. Clin Chem, 1999. 45(11): p. 1960-6). Nam and associates (Nam, R.K., et al., Serum human glandular kallikrein-2 protease levels predict the presence of prostate cancer among men with elevated prostate-specific antigen. J Clin Oncol, 2000. 18(5): p. 1036-42) documented that serum human glandular kallikrein 2 levels predict the presence of prostate cancer among men with elevated PSA.

Ki-67

[0520] This is a nuclear protein that is expressed in proliferating normal and neoplastic cells. Ki-67 expression occurs during the phase of the cell cycle designated as late G1, S, M, and G2, but not in G0 phase (Cattoretti, G., et al., Monoclonal antibodies against recombinant parts of the Ki-67 antigen (MIB 1 and MIB 3) detect proliferating cells in microwave-processed formalin-fixed paraffin sections. J Pathol, 1992. 168(4): p. 357-63). Ki-67 is commonly used as a cell proliferation marker and it is located in the nucleus. Studies show that Ki-67 staining in prostate cancer provides independent prognostic information after radical prostatectomy and Ki-67 immunoreactivity is a predictor for prostate cancer survival (Halvorsen, O.J., et al., Maximum Ki-67 staining in prostate cancer provides independent prognostic information after radical prostatectomy. Anticancer Res, 2001. 21(6A): p. 4071-6; Stattin, P., et al., Cell proliferation

assessed by Ki-67 immunoreactivity on formalin fixed tissues is a predictive factor for survival in prostate cancer. J Urol, 1997. 157(1): p. 219-22).

Leu 7

[0521] This antigen is present on non-cancerous and cancerous prostatic epithelia, as well as natural killer cells and myelinated nerves. Liu and associates (Liu, X., et al., Immunohistochemical study of HNK-1 (Leu-7) antigen in prostate cancer and its clinical significance. Chin Med J (Engl), 1995. 108(7): p. 516-21) showed that 94% of prostate cancer is positive for Leu 7. Well-differentiated cancer showed the highest percentage of positive cancer cells and the strongest staining, while poorly differentiated cancer had the lowest percentage of positive cancer cells and the weakest staining. Another study suggested that the expression of Leu 7 on prostate cancer may be a useful prognostic factor for patients with prostate cancer (Liu, X.H., et al., The prognostic value of the HNK-1 (Leu-7) antigen in prostatic cancer--an immunohistochemical study. Hinyokika Kiyo, 1993. 39(5): p. 439-44).

MDM2

[0522] This is a cellular proto-oncogene product. MDM2 binds to p53, promoting degradation via ubiquitin, masking its transactivation domain, and inhibiting its transcriptional activation of genes related to cell cycle arrest and apoptosis (Momand, J. and G.P. Zambetti, Mdm-2: "big brother" of p53. J Cell Biochem, 1997. 64(3): p. 343-52). Amplification of the MDM2 gene or over-expression of the MDM2 protein have been implicated in the development of tumors, and MDM2 over-expression has been related to more aggressive disease and poorer survival (Freedman, D.A., L. Wu, and A.J. Levine, Functions of the MDM2 oncoprotein. Cell Mol Life Sci, 1999. 55(1): p. 96-107). Leite and associates (Leite, K.R., et al., Abnormal expression of MDM2 in prostate carcinoma. Mod Pathol, 2001. 14(5): p. 428-36) showed MDM2 was over-expressed in 41% of prostate adenocarcinomas cases. Tumors that were positive for both p53 and MDM2 were larger and of more advanced stage. Results suggest that MDM2-positive/p53-positive phenotype identifies prostate cancers with aggressive behavior.

N-Cadherin

[0523] Similar to E-Cadherin, this is a cell adhesion molecule. Changes in cell-cell interactions are critical in the process of cancer progression. Likewise, it has been shown that loss of expression of the cell adhesion molecule E-cadherin is associated with grade, stage, and prognosis in many carcinomas, including prostate cancer. Impaired E-cadherin-mediated

interactions result in an invasive phenotype. However, the mere loss of cell-cell contact and communication is not the sole explanation for the observed correlation between loss of Ecadherin-mediated adhesion and poor clinical outcome (Kuniyasu, H., et al., Relative expression of type IV collagenase, E-cadherin, and vascular endothelial growth factor/vascular permeability factor in prostatectomy specimens distinguishes organ-confined from pathologically advanced prostate cancers. Clin Cancer Res, 2000. 6(6): p. 2295-308; Bryden, A.A., et al., E-cadherin and beta-catenin are down-regulated in prostatic bone metastases. BJU Int, 2002. 89(4): p. 400-3; Umbas, R., et al., Decreased E-cadherin expression is associated with poor prognosis in patients with prostate cancer. Cancer Res, 1994. 54(14): p. 3929-33). Bussemakers and associates (Bussemakers, M.J., et al., Complex cadherin expression in human prostate cancer cells. Int J Cancer, 2000. 85(3): p. 446-50) demonstrated that N-Cadherin is up-regulated in human prostate cancer cell lines. Another study showed that N-cadherin was not expressed in normal prostate tissue, however, in prostatic cancer N-cadherin was found to be expressed in the poorly differentiated areas, which showed mainly aberrant or negative E-cadherin staining (Tomita, K., et al., Cadherin switching in human prostate cancer progression. Cancer Res, 2000. 60(13): p. 3650-4).

P504S

[0524] Alpha-methylacyl-CoA racemase (P504S) is a cytoplasmic protein. It was recently identified by cDNA library subtraction in conjunction with high throughput microarray screening from prostate carcinoma. Jiang and associates (Jiang, Z., et al., P504S: a new molecular marker for the detection of prostate carcinoma. Am J Surg Pathol, 2001. 25(11): p. 1397-404) examined P504S by immunocytochemistry on benign and cancerous prostate tissues. P504S showed strong cytoplasmic granular staining in 100% of the prostate carcinomas regardless of Gleason scores and diffuse. In contrast, 171 of 194 (88%) of benign prostates, including 56 of 67 (84%) benign prostate cases and 115 of 127 (91%) cases of benign glands adjacent to cancers, were negative for P504S. Another study revealed P504S is a very useful marker for distinguishing the atypical adenomatous hyperplasia of the prostate from prostatic adenocarcinoma (Yang, X.J., et al., Expression of alpha-Methylacyl-CoA racemase (P504S) in atypical adenomatous hyperplasia of the prostate. Am J Surg Pathol, 2002. 26(7): p. 921-5).

p53

[0525] This is a tumor suppressor gene. Inactivation of p53 is implicated in tumorigenesis for over half of all human cancers. It functions as a transcriptional regulator involved in G₁ phase growth arrest of cells in response to DNA damage, as well as having a role in the regulation of the spindle checkpoint, centrosome homeostasis, and G₂-M phase transition. It also induces apoptosis by transcription-dependent and independent mechanisms in many cell types and regulates tumor angiogenesis (Kirsch, D.G. and M.B. Kastan, Tumor-suppressor p53: implications for tumor development and prognosis. J Clin Oncol, 1998. 16(9): p. 3158-68; Agarwal, M.L., et al., The p53 network. J Biol Chem, 1998. 273(1): p. 1-4; Liebermann, D.A., B. Hoffman, and R.A. Steinman, Molecular controls of growth arrest and apoptosis: p53-dependent and independent pathways. Oncogene, 1995. 11(1): p. 199-210). Thompson and associates (Thompson, S.J., et al., P53 and Ki-67 immunoreactivity in human prostate cancer and benign hyperplasia. Br J Urol, 1992. 69(6): p. 609-13) showed prostate cancer specimens with p53 were stained, whereas no staining was observed in benign prostate hyperplasia (BPH). Another study showed that nuclear accumulation of p53 was a significant prognostic indicator for prostate cancer (Quinn, D.I., et al., Prognostic significance of p53 nuclear accumulation in localized prostate cancer treated with radical prostatectomy. Cancer Res, 2000. 60(6): p. 1585-94).

Prostate Acid Phosphatase (PAP)

[0526] Prostate acid phosphatase (PAP), as prostate specific antigen (PSA), has greatly increased the feasibility of reliable diagnosis of primary or metastatic prostatic carcinoma. Studies show that diagnostic sensitivity of PAP is 90-100% and its specificity is 87-100%, and PAP is a useful prognostic indicator of advanced prostatic carcinoma (Svanholm, H. and M. Horder, Clinical application of prostatic markers. I. Classification of prostatic tumours using immunohistochemical techniques. Scand J Urol Nephrol Suppl, 1988. 107: p. 65-70; (Sakai, H., et al., Prostate specific antigen and prostatic acid phosphatase immunoreactivity as prognostic indicators of advanced prostatic carcinoma. J Urol, 1993. 149(5): p. 1020-3).

Prostate Inhibin Peptide (PIP)

[0527] Prostate inhibin peptide (PIP) is a polypeptide synthesized by the prostate gland. It is involved in prostatic growth and differentiation. The PIP gene is localized in the 7q34 region that contains a number of fragile sites. Rearrangement of PIP genes was found in prostate carcinomas (Autiero, M., et al., Abnormal restriction pattern of PIP gene associated with human

primary prostate cancers. DNA Cell Biol, 1999. 18(6): p. 481-7). Garde and associates (Garde, S.V., et al., Prostate inhibin peptide (PIP) in prostate cancer: a comparative immunohistochemical study with prostate-specific antigen (PSA) and prostatic acid phosphatase (PAP). Cancer Lett, 1994. 78(1-3): p. 11-7) showed PIP is as sensitive and specific of an immunohistochemical marker as PSA and PAP in the diagnosis of prostate carcinoma. Further, the androgen-independent nature of PIP may give it an advantage over PSA/PAP in tumors exposed to androgen ablating agents.

Prostate Specific Antigen (PSA)

[0528] Prostate specific antigen (PSA) is a product of prostatic epithelium and is normally secreted in the semen. It is a serine protease whose function is to cleave and liquefy the seminal coagulum formed after ejaculation. PSA is organ specific, not cancer specific. Thus, elevations in PSA levels occur not only in cancer, but also in non-neoplastic conditions, such as nodular hyperplasia of the prostate and prostatitis. PSA by itself cannot be used for detection of early cancer. When combined with a rectal examination and transrectal ultrasonography, however, measurement of PSA levels is considered useful in detection of early-stage cancers (Crawford, J.M. and R.S. Cotran, The Male Genital Tract, in Robbins Pathologic Basis of Disease, R.S. Cotran, V. Kuman, and T. Collins, Editors. 1999, W. B. Saunders Company: Philadelphia. p. 1011-1034). More evidence suggests that free serum PSA is more accurate than total PSA in the diagnosis of prostate carcinoma (Catalona, W.J., et al., Use of the percentage of free prostatespecific antigen to enhance differentiation of prostate cancer from benign prostatic disease: a prospective multicenter clinical trial. Jama, 1998. 279(19): p. 1542-7). Immunohistochemical study of PSA for prostate carcinoma diagnosis is also very sensitive and specific. Svanholm and associates (Svanholm, H. and M. Horder, Clinical application of prostatic markers. I. Classification of prostatic tumours using immunohistochemical techniques. Scand J Urol Nephrol Suppl, 1988. 107: p. 65-70) demonstrated the sensitivity of PSA for prostate carcinoma diagnosis is 94-100% and its specificity is 100%.

V. Breast Cancer

[0529] Non-invasive mammography as a screening tool for breast cancer is not effective (Gotzsche, P.C. and O. Olsen, *Is screening for breast cancer with mammography justifiable?*Lancet, 2000. 355(9198): p. 129-34). Therefore, other techniques for screening for breast cancer have been studied. A variety of breast cancer markers have been discovered to aid physicians in

making timely, precise diagnoses, and to provide significantly better patient management. Unfortunately, none of these tumor makers is a "magic bullet" with both high sensitivity and specificity. Therefore, alternative ways to enhance diagnostic accuracy are necessary. [0530] An alternative way to enhance diagnostic accuracy is to develop a panel comprising a plurality of probes each of which specifically binds a marker associated with breast cancer. All candidate probes are to be tested with ICC and/or IHC techniques. In some embodiments, specimens may be obtained from fine needle aspirations (FNA). In other embodiments, specimens may be obtained from fine needle biopsies (FNB). Proper location and sampling of tumors by FNA and FNB procedures may be aided by ultrasound and other image-guiding techniques. In some embodiments, test cells are obtained from breast ductal lavage. Devices for this purpose are readily available and may work by collecting cells through an aspirator similar to a manual breast pump. A small suction cup is used to draw nipple aspirate fluid (NAF) through the nipple. The presence of NAF helps locate natural openings of the ducts on the surface of the nipple. Then a tiny, flexible microcatheter is inserted approximately half an inch into the duct to be lavaged in order to collect cells lining the breast duct.

Statistical analysis will be used to design panels, as described above for lung cancer. During processing, technical issues such as cell smears or pellets not sticking to slides during harsh washings may occur in some embodiments. However, such issues can readily be addressed by manipulation of software or modifying staining protocols to mitigate such problems. In some embodiments, the specimens will be processed and analyzed using a device that automatically samples the specimen and prepares slides for diagnosis. It is anticipated that a broad menu of probes will be used initially. The number of probes will be pruned to a suitably sized panel in order to retain a high level of sensitivity and specificity. Selection of the final probes will be based on a pre-defined threshold of the percentage of positive stained tumor cells. Sophisticated statistical analysis will be employed to make these determinations. Since the panel-assay approach to detecting malignancies is applicable to solid tumors, and several of the same tumor markers are in different panels, this method may be carried out in parallel, as well as serially. In this manner, the assay development process can be expedited.

Library of Probes/Markers

[0532] Various sources containing information about cancer markers were reviewed. An arbitrary criterion of 20% or greater positivity of breast cancer was used to select probes for a preferred panel for detection and/or diagnosis of breast cancer. The term "20% or greater positivity" means that if 100 tumor cases were studied, 20 or more of these cases would have shown a presence of the individual marker, while the remaining 80 cases would not have shown a presence of the individual marker. A preferred panel may include molecular markers selected from AE1/AE3, BCA-225, Bcl-2, BRCA-1, Cancer Antigen 15.3 (CA 15.3), Cathespin D, Carcinoembryonic Antigen (CEA), C-erb-B2, E-Cadherin, Epidermal Growth Factor Receptor (EGFR), Estrogen receptor (ER), Gross Cystic Disease Fluid Protein 15 (GCDFP-15), HOX-B3, Ki-67, MUC-1, p53, p65, Progesterone Receptor (PR), Retinoblastoma (Rb) and Transglutaminase K (TGK). A brief description of the library of probes/markers utilized in the present example is provided below.

AE1/AE3

[0533] This is a cocktail of anti-keratin antibodies. Keratins are a group of water-insoluble proteins that form an intermediate filament in the cells of epithelial origin, such as breast ductal epithelium. Anti-keratin AE1 recognizes the 56 and 40 kD keratins of the acidic sub-family. Anti-keratin AE3 recognizes the basic sub-family. AE1 and AE3 have been shown to be more effective than other anti-cytokeratins in identification of lymph node metastasis from breast ductal carcinoma (Elson, C.E., D. Kufe, and W.W. Johnston, Immunohistochemical detection and significance of axillary lymph node micrometastases in breast carcinoma. A study of 97 cases. Anal Quant Cytol Histol, 1993. 15(3): p. 171-8). Kowolik and associates (Kowolik, J.H., et al., Detection of micrometastases in sentinel lymph nodes of the breast applying monoclonal antibodies AE1/AE3 to pancytokeratins. Oncol Rep, 2000. 7(4): p. 745-9) showed 32 of 33 cases of axillary lymph nodes with breast cancer metastasis that were correctly predicted by AE1/AE3 immunohistochemical staining. AE1/AE3 has also been shown to be the most sensitive marker for detecting occult metastasis in nodes of infiltrating lobular breast carcinoma (Kainz, C., et al., Infiltrating lobular breast carcinoma: detection of occult regional lymph node metastasis by immunohistochemistry. Anticancer Res, 1993. 13(1): p. 73-4).

BCA-225

[0534] This antibody recognizes a human breast carcinoma associated glycoprotein BCA-225 (220-225 kD), which differs in size and distribution from other breast carcinoma antigens. However, unlike other carcinoma antibodies against breast carcinoma antigens, BCA-225 does not react with benign or malignant gastrointestinal tissues. One study showed 94% of breast carcinoma are positive for BCA-225 (Mesa-Tejada, R., et al., *Immunocytochemical distribution of a breast carcinoma associated glycoprotein identified by monoclonal antibodies*. Am J Pathol, 1988. 130(2): p. 305-14), while another showed 78% of effusion containing breast carcinoma was positive and all benign effusions were negative for BCA-225 by immunocytochemistry. BCA-225 is highly specific discriminator and very useful in differential diagnosis of adenocarcinoma and reactive mesothelial cells (Loy, T.S., A.A. Diaz-Arias, and J.T. Bickel, *Value of BCA-225 in the cytologic diagnosis of malignant effusions: an immunocytochemical study of 197 cases.* Mod Pathol, 1990. 3(3): p. 294-7).

Bcl-2

[0535] Survival threshold for a cell is determined by the balance between cell-death suppressor and cell-death promoter signals provided by external factors or stimuli, as well as by intracellular molecules. Bcl-2 has a central role in this determination and its product acts as an anti-apoptotic molecule (Daidone, M.G., et al., Clinical studies of Bcl-2 and treatment benefit in breast cancer patients. Endocr Relat Cancer, 1999. 6(1): p. 61-8). The Bcl-2 protein has been shown to contribute to oncogenesis because it can transform and immortalize cells in cooperation with c-myc, ras, or viral genes (Del Bufalo, D., et al., Bcl-2 overexpression enhances the metastatic potential of a human breast cancer line. Faseb J, 1997. 11(12): p. 947-53). Bcl-2 expression is most commonly associated with the t (14; 18) translocation in most follicular lymphomas. More recently, Bcl-2 has been identified in non-hematologic malignancies. Alsabeh and associates determined that Bcl-2 is a useful marker in distinguishing metastatic breast carcinoma from primary lung and gastric carcinomas, and it is a useful prognostic indicator as well (Alsabeh, R., et al., Expression of bcl-2 by breast cancer: a possible diagnostic application. Mod Pathol, 1996. 9(4): p. 439-44).

BRCA-1

[0536] This is a tumor suppressor gene located on the long arm of chromosome 17. Tumor suppressor genes play a critical role in regulating cell growth. BRCA-1 is a nuclear

phosphoprotein, which normally functions as a negative regulator of the cell cycle and may be an active inhibitor of neoplastic progression. Mutation of the BRCA1 gene has been demonstrated in 80% of familial breast cancer. Decreased mRNA levels or aberrant sub-cellular locations of BRCA1 have been identified in breast cancer lines and in sporadic cases of breast cancer tissues. BRCA1 mutations are linked to ovarian cancer as well (Lee, W.Y., et al., Immunolocalization of BRCA1 protein in normal breast tissue and sporadic invasive ductal carcinomas: a correlation with other biological parameters. Histopathology, 1999. 34(2): p. 106-12). Jarvis and associates (Jarvis, E.M., J.A. Kirk, and C.L. Clarke, Loss of nuclear BRCA1 expression in breast cancers is associated with a highly proliferative tumor phenotype. Cancer Genet Cytogenet, 1998. 101(2): p. 109-15) found that nuclear staining for BRCA-1 was observed in most sporadic tumors, but nuclear BRCA-1 was reduced or absent in the majority of familial and early onset breast tumors. A significant inverse correlation was found between nuclear BRCA-1 and expression of the proliferation marker Ki-67. Another study revealed BRCA-1 expression was correlated with other prognostic markers including p53, c-erb-B2, Bcl-2, ER, histological grade, tumor size, axillary lymph node status and age (Lee, W.Y., et al., Immunolocalization of BRCA1 protein in normal breast tissue and sporadic invasive ductal carcinomas: a correlation with other biological parameters. Histopathology, 1999. 34(2): p. 106-12).

Cancer Antigen 15.3 (CA-15.3)

[0537] Cancer antigen 15.3 is a serum carbohydrate antigen. Increased serum concentration of CA-15.3 has been associated with breast carcinoma as well as normal and benign breast disease. However, the level of CA-15.3 is significantly higher in breast carcinoma than that in either normal or benign breast disease (Barak, M., et al., CA-15.3, TPA and MCA as markers for breast cancer. Eur J Cancer, 1990. 26(5): p. 577-80). Martoni and associates (Martoni, A., et al., CEA, MCA, CA 15.3 and CA 549 and their combinations in expressing and monitoring metastatic breast cancer: a prospective comparative study. Eur J Cancer, 1995. 31A(10): p. 1615-21) have documented that CA 15.3 has higher sensitivity than other tumor markers in detecting metastatic breast cancer.

Cathepsin D

[0538] This is a soluble lysosomal aspartic proteinase. It is synthesized in the endoplasmic reticulum as a preprocathepsin D. Having a mannose-6-phosphate tag, procathepsin D is recognized by a mannose-6-phosphate receptor. Upon entering into an acidic lysosome, single-

chain procathepsin D (52 kD) is activated to cathepsin D. The fundamental role of cathepsin D is the degradation of intracellular and internalized proteins. Increased levels of cathepsin D (both at the mRNA and protein levels) were first reported in several human neoplastic tissues in the mideighties. These findings generated intense research in a possible role for cathepsin D in neoplastic processes. A strong predictive value was found for cathepsin D concentrations in breast cancer, as well as many other tumor types (Vetvicka, V., et al., Analysis of the interaction of procathepsin D activation peptide with breast cancer cells. Int J Cancer, 1997. 73(3): p. 403-9; Vetvicka, V., J. Vetvickova, and M. Fusek, Effect of procathepsin D and its activation peptide on prostate cancer cells. Cancer Lett, 1998. 129(1): p. 55-9). Niu and associates (Niu, Y., et al., Potential markers predicting distant metastasis in axillary node-negative breast carcinoma. Int J Cancer, 2002. 98(5): p. 754-60) found Cathepsin D to be a potential marker predicting distant metastasis in axillary node-negative breast cancer patients.

Carcinoembryonic Antigen (CEA)

[0539] Carcinoembryonic antigen (CEA) is well known for its role in diagnosis and follow-up for colorectal cancer. CEA positivity of breast carcinoma is also reported. Alexiev (Alexiev, B.A., et al., Immunocytochemical detection of carcinoembryonic antigen in fine-needle aspirates from patients with diverse breast diseases. Diagn Cytopathol, 1993. 9(4): p. 377-82) and associates reported that 90% of primary breast carcinomas showed positive cytoplasmic staining for CEA whereas none of the benign breast diseases are positive on fine-needle aspirates and paraffin-embedded tissue sections. Others have reported CEA was positive for fibroadenoma and cystic change disease ranging from 25% to 64% (Wittekind, C., S. Von Kleist, and W. Sandritter, CEA positivity in tissue and sera of patients with benign breast lesions. Oncodev Biol Med, 1981. 2(6): p. 381-90).

C-erb-B2

[0540] This is an oncogene and its product is known as oncoprotein. C-erb-B2, also known as Her2/neu, belongs to the epidermal growth factor receptor (EGFR) family that includes c-erb-B1, c-erb-B2, and c-erb-B3. Over-expression of c-erb-B2 is associated with a high percentage of human carcinomas arising within the breast, ovary, lung, prostate, stomach and salivary glands. It is suggested that tumors that over-express the growth factors, such as c-erb-B2, would be exquisitely sensitive to the growth-promoting effects of a small amount of growth factors and hence likely to be more aggressive. This hypothesis is supported by the observation that high

levels of c-erb-B2 protein on breast cancer cells are a harbinger of poor prognosis (Mitchell, R.N. and R.S. Cotran, Neoplasia, in Robbins Pathologic Basis of Disease, R.S. Cotran, V.K. Kumar, and T. Collins, Editors. 1999, W. B. Sauders Company: Philadelphia. p. 260-327). Inaji and associates (Inaji, H., et al., ErbB-2 protein levels in nipple discharge: role in diagnosis of early breast cancer. Tumour Biol, 1993. 14(5): p. 271-8) found that levels of c-erb-B2 oncoprotein in nipple discharge were elevated in breast carcinoma patients. One study revealed that 34% of fine needle aspirations of breast carcinoma and all but one corresponding tissue section were positive for c-erb-B2 oncoprotein (Jorda, M., P. Ganjei, and M. Nadji, Retrospective c-erbB-2 immunostaining in aspiration cytology of breast cancer. Diagn Cytopathol, 1994. 11(3): p. 262-5).

E-Cadherin

[0541] This protein is suggested to be the major cell adhesion molecule in mammary glands. In cytoplasm, E-Cadherin is linked to alpha- and beta-catenin which mediates the connection of the cytoskeleton. In addition, c-erbB-2 oncoprotein causes disruption of the cell adhesion system through beta-catenin phosphorylation (Nagae, Y., et al., Expression of E-cadherin catenin and C-erbB-2 gene products in invasive ductal-type breast carcinomas. J Nippon Med Sch, 2002. 69(2): p. 165-71). Decreased expression of E-Cadherin is found in breast carcinoma (Mitchell, R.N. and R.S. Cotran, Neoplasia, in Robbins Pathologic Basis of Disease, R.S. Cotran, V.K. Kumar, and T. Collins, Editors. 1999, W. B. Sauders Company: Philadelphia. p. 260-327).

Epidermal Growth Factor Receptor (EGFR)

[0542] Epidermal growth factor receptor (EGFR) is a transmembrane glycoprotein. It plays an important role in cell growth and differentiation. EGFR expression has been shown in a broad spectrum of normal tissues, whereas over-expression has been associated with a variety of neoplasms. Sue and associates (Suo, Z., et al., Type 1 protein tyrosine kinases in benign and malignant breast lesions. Histopathology, 1998. 33(6): p. 514-21) examined the expression pattern of the four EGFR family members in breast tumor tissues and found 53% of breast tumor tissues were strongly positive for EGFR, though benign tumors also expressed EGFR protein but all at a lower, moderate level. An association between EGFR expression and increasing malignancy grade was found in a group of infiltrating ductal carcinomas.

Estrogen Receptor (ER)

[0543] Estrogens control a variety of physiological and disease-linked processes, most notably reproduction, bone remodeling and breast cancer, and their effects are transduced through classic receptors referred to as estrogen receptor (ER). This monoclonal antibody reacts with the N-terminal domain (A/B region) of the 67 kD polypeptide chain of the estrogen receptor, and exhibits a nuclear staining pattern with little or no cytoplasmic reactivity. In tumor tissue, ER reacts strongly with epithelial cells of breast cancers and human neoplasms derived from other estrogen-dependent tissues. In general, cancers that have cells expressing ER in their nuclei will have better prognoses because such positive neoplastic cells are better differentiated and can respond to hormonal manipulation. Tamoxifen, a drug, is often utilized for this purpose (Barnes, D.M., et al., *Immunohistochemical determination of oestrogen receptor: comparison of different methods of assessment of staining and correlation with clinical outcome of breast cancer patients.* Br J Cancer, 1996. 74(9): p. 1445-51; Pichon, M.F., et al., *Prognostic value of steroid receptors after long-term follow-up of 2257 operable breast cancers.* Br J Cancer, 1996. 73(12): p. 1545-51).

Gross Cystic Disease Fluid Protein 15 (GCDFP-15)

[0544] Gross Cystic Disease Fluid Protein 15 (GCDFP-15) is a glycoprotein (15 kD) expressed by apocrine sweat glands, eccrine glands, minor salivary glands, bronchial glands and metaplastic epithelium of the breast. Breast carcinomas (primary and metastatic lesions) with apocrine features express the GCDFP-15 antigen. GCDFP-15 is positive in extra-mammary Paget's disease, while other tumors tested negative. Numerous histopathologic studies have shown GCFDP-15 to be a specific marker for breast cancer in surgical specimens. Fiel and associates (Fiel, M.I., et al., *Value of GCDFP-15 (BRST-2) as a specific immunocytochemical marker for breast carcinoma in cytologic specimens*. Acta Cytol, 1996. 40(4): p. 637-41) also demonstrated that GCFDP-15 is a specific immunocytochemical marker for breast carcinoma in cytologic specimens.

HOX-B3

[0545] The homeobox (HOX) genes encode proteins which contain 61 amino acid DNA-binding homeodomain and are involved in the transcriptional regulation of other genes during normal onco- and histogenesis. Class I HOX genes are organized into four clusters on different chromosomes in humans, with a high conservation in the order of the genes within each of these

clusters. Re-expression of HOX gene products has been reported in a wide variety of neoplastically transformed cells and it seems quite likely that the HOX genes represent yet another class of oncofetal antigens involved in normal development and carcinogenesis, as well as tumor progression. HOX-3 is one of the HOX gene products (HOX-B3, -B4, and -C6). One study showed over 90% of breast carcinoma is positive for HOX-B3 (Bodey, B., et al., Immunocytochemical detection of the homeobox B3, B4, and C6 gene products in breast carcinomas. Anticancer Res, 2000. 20(5A): p. 3281-6).

Ki-67

[0546] This is a nuclear protein that is expressed in proliferating normal and neoplastic cells. Ki-67 expression occurs during the phase of the cell cycle designated as late G1, S, M, and G2. However during the G0 phase, the antigen cannot be detected. Studies have shown that expression of Ki-67 was inversely associated with estrogen and progesterone receptors, suggesting that a high Ki-67 level seems to characterize a more aggressive phenotype and poor prognosis (Ceccarelli, C., et al., Quantitative p21(waf-1)/p53 immunohistochemical analysis defines groups of primary invasive breast carcinomas with different prognostic indicators. Int J Cancer, 2001. 95(2): p. 128-34; Ceccarelli, C., et al., Retinoblastoma (RB1) gene product expression in breast carcinoma. Correlation with Ki-67 growth fraction and biopathological profile. J Clin Pathol, 1998. 51(11): p. 818-24).

MUC-1

[0547] Epithelial mucins are glycoproteins secreted by epithelial cells and their carcinomas. At least nine mucin genes have been identified, and their products, MUC1 thru MUC9, are expressed in various epithelia. MUC1 is a mucin expressed in breast epithelial cells, whereas MUC2 and MUC3 are primarily intestinal mucins. MUC-1 antigen is a cell surface glycoprotein. This antigen is abundant in 90% of human breast cancers in forms not present in normal tissue (Diaz, L.K., E.L. Wiley, and M. Morrow, Expression of epithelial mucins Muc1, Muc2, and Muc3 in ductal carcinoma in situ of the breast. Breast J, 2001. 7(1): p. 40-5). One study (Croce, M.V., et al., Expression of tumour associated antigens in normal, benign and malignant human mammary epithelial tissue: a comparative immunohistochemical study. Anticancer Res, 1997. 17(6D): p. 4287-92) showed benign breast tissues expressed a low intensity of MUC-1, restricted to apical cell surface membranes and lumen debris.

p53

[0548] This is tumor suppressor gene. High concentrations of the p53 protein occur in a large number of tumors and tumor cell lines, while it is present in only minute amounts in normal cells and tissues. The increased expression in tumor cells may be caused by mutation of the p53 protein or by complexing with other proteins. The gene for p53 is located on chromosome 17p, a frequent site of allele loss in tumors of the breast, lung, colon, ovaries, testicles, bladder, brain, melanomas, certain types of leukemia and neurofibrosarcoma. Tumor cell expression of p53 protein can be detected by immuno-histochemistry, exhibiting a nuclear staining pattern. Expression of the p53 oncoprotein has been shown to correlate with poor prognosis in breast cancer (Lee, W.Y., et al., Immunolocalization of BRCA1 protein in normal breast tissue and sporadic invasive ductal carcinomas: a correlation with other biological parameters. Histopathology, 1999. 34(2): p. 106-12; Midulla, C., et al., Immunohistochemical expression of p53, nm23-HI, Ki67 and DNA ploidy: correlation with lymph node status and other clinical pathologic parameters in breast cancer. Anticancer Res, 1999. 19(5B): p. 4033-7).

p65

[0549] This 65 kD oncofetal protein has been identified as a new member of the steroid/thyroid super-family of genes, with as yet an unknown ligand. It is suggested that this receptor may play an important role in the development of tumors. The altered form of p65 is linked to the overproduction of certain hormones that may cause breast cancers (Hanausek, M., et al., *The oncofetal protein p65: a new member of the steroid/thyroid receptor superfamily.* Cancer Detect Prev, 1996. 20(2): p. 94-102). Mirowski and associates (Mirowski, M., et al., *Serological and immunohistochemical detection of a 65-kDa oncofetal protein in breast cancer.* Eur J Cancer, 1994. 30A(8): p. 1108-13) revealed that p65 was positive in 90% of sera from breast cancer patients and positive in 80% of corresponding biopsied tissue assessed by immunohistochemistry. Results indicate p65 may be a potential serum and/or immunohistochemical marker for breast cancer.

Progesterone Receptor (PR)

[0550] The monoclonal antibody to progesterone receptor (PR) exhibits a nuclear staining pattern. In tumor tissue, PR expression is strongly in epithelial cells of breast cancers and human neoplasms derived from other progesterone-dependent tissues, while in normal tissues it is positive in mammary glands and the uterus. The significance of PR positivity in a breast

carcinoma is less well understood. In general, cancers that are ER positive will also be PR positive. However, carcinomas that are PR positive, but not ER positive, may have a worse prognosis (Blanco, G., et al., Estrogen and progesterone receptors in breast cancer: relationships to tumour histopathology and survival of patients. Anticancer Res, 1984. 4(6): p. 383-9). One study showed PR was positive in all cutaneous metastatic breast tumors whereas only one tumor was positive for ER (Wallace, M.L. and B.R. Smoller, Differential sensitivity of estrogen/progesterone receptors and BRST-2 markers in metastatic ductal and lobular breast carcinoma to the skin. Am J Dermatopathol, 1996. 18(3): p. 241-7).

Retinoblastoma (Rb)

[0551] This is one of the tumor suppressor genes. Retinoblastoma protein (pRb) is a protein that is encoded by the retinoblastoma gene and functions to regulate the cell cycle at G0/G1. Loss of Rb function leads to uncontrolled cell growth. Inactivation of the retinoblastoma gene is documented in various types of cancer, including breast cancer. Retinoblastoma gene underexpression promotes breast-tumor aggressiveness and rapid tumor-cell proliferation (Bieche, I. and R. Lidereau, Loss of heterozygosity at 13q14 correlates with RB1 gene underexpression in human breast cancer. Mol Carcinog, 2000. 29(3): p. 151-8). Ceccarelli and associates (Ceccarelli, C., et al., Retinoblastoma (RB1) gene product expression in breast carcinoma. Correlation with Ki-67 growth fraction and biopathological profile. J Clin Pathol, 1998. 51(11): p. 818-24) studied pRb expression and tumor markers Ki-67, ER/PR, p53 and EGFR in invasive breast carcinoma and found that pRb expression paralleled proliferative activity in a majority of breast carcinomas examined, suggesting that in these cases the protein behaves normally in regulating the cell cycle. Conversely, in cases with a loss of pRb immunostaining, the combined expression of specific highly aggressive factors, such as EGFR and p53 expression, estrogen receptor/progesterone receptor negative status and high K67, seems to characterize a more aggressive phenotype.

Transglutaminase K (TGK)

[0552] Transglutaminase K, is not well studied for breast cancer diagnosis. However, Friedrich and associates (Friedrich, M., et al., Correlation between immunoreactivity for transglutaminase K and for markers of proliferation and differentiation in normal breast tissue and breast carcinomas. Eur J Gynaecol Oncol, 1998. 19(5): p. 444-8) showed weak to strong membrane staining was detected in 17 of 30 breast carcinomas, while 90% of normal breast tissue revealed

no immunoreactivity to TGK. The results suggest that up-regulation of TGK in breast carcinomas may play an important role in the regulation of tumor cell invasive properties by modulating cell-matrix interactions or by facilitating the assembly of matrix and tissue remodeling.

VI. Cervical Cancer

[0553] A variety of cervical cancer markers have been discovered to aid physicians in making timely, precise diagnoses, and to provide significantly better patient management. Unfortunately, none of these tumor makers is a "magic bullet" with both high sensitivity and specificity. Therefore, alternative ways to enhance diagnostic accuracy are necessary.

[0554] An alternative way to enhance diagnostic accuracy is to develop a panel comprising a plurality of probes each of which specifically binds a marker associated with cervical cancer. All candidate probes are to be tested with ICC and or IHC techniques. In some embodiments, specimens may be obtained using Pap smears. Conventional Pap smears utilize spatulas and brushes to collect cervical cells. For the liquid-based preparation (LBP) Pap tests, cells may be collected using a broom, brush, or balloon. For example, Cytyc's ThinPrep® product and a TriPath's SurePrepTM product may be used. Additionally Molecular Diagnostics' "e2 CollectorTM" may be used for obtaining cervical cells. It is a silicone balloon, shaped like a mirror image of the cervix. When inflated against the cervix, cells adhere to the balloon's surface and collect endocervix and ectocervix cells in a single step.

[0555] Once the specimens are collected, the specimens will be processed and analyzed. Statistical analysis will be used to design panels, as described above for lung cancer. During processing, technical issues such as cell smears or pellets not sticking to slides during harsh washings may occur in some embodiments. However, such issues can readily be addressed by manipulation of software or modifying staining protocols to mitigate such problems. In some embodiments, the specimens will be processed and analyzed using a device that automatically samples the specimen and prepares slides for diagnosis. It is anticipated that a broad menu of probes will be used initially. The number of probes will be pruned to a suitably sized panel in order to retain a high level of sensitivity and specificity. Selection of the final probes will be based on a pre-defined threshold of the percentage of positive stained tumor cells. Sophisticated statistical analysis will be employed to make these determinations. Since the panel-assay approach to detecting malignancies is applicable to solid tumors, and several of the same tumor

markers are in different panels, this method may be carried out in parallel, as well as serially. In this manner, the assay development process can be expedited.

[0556] Epidemiologic studies suggest that carcinoma of the cervix is caused by a sexually transmitted agent, and human papillomavirus (HPV) is a prime suspect. Approximately 70 genetically distinct types of HPV have been identified. Types 16 and 18, less commonly, types 31, 33, 35, and 51 are found in approximately 85% of invasive squamous cell cancers and their precursors (severe dysplasias and carcinoma in situ). In contrast to cervical cancers, genital warts with low malignant potential are associated with "low risk" types of HPV-6 and HPV-11. (Mitchell, R.N. and R.S. Cotran, Neoplasia, in Robbins Pathologic Basis of Disease, R.S. Cotran, V. Kumar, and T. Collins, Editors. 1999, W. B. Saunders Company: Philadelphia. p. 260-327). [0557] Molecular diagnosis of HPV infection is more sensitive and specific than conventional Pap smears, and has added value in the evaluation of women with equivocal Pap smear results. Digene's Hybrid Capture II HPV DNA test is highly effective in detecting patients with highgrade dysplasia. (Solomon, D., M. Schiffman, and R. Tarone, Comparison of three management strategies for patients with atypical squamous cells of undetermined significance: baseline results from a randomized trial. J Natl Cancer Inst, 2001. 93(4): p. 293-9). Another HPV method is detection of E6 and E7 proteins of HPV 16 and HPV 18 developed by Molecular Diagnostics. The oncogenic potential of HPV-16 and HPV-18 has been related to these two early viral gene products: (Huibregtse, J.M. and S.L. Beaudenon, Mechanism of HPV E6 proteins in cellular transformation. Semin Cancer Biol, 1996. 7(6): p. 317-26) and (zur Hausen, H., Papillomavirus and p53. Nature, 1998. 393(6682): p. 217). Roche Diagnostics recently acquired HPV testing patents from Institut Pasteur.

Library of Probes/Markers

[0558] Various sources containing information about cancer markers were reviewed. An arbitrary criterion of 20% or greater positivity of breast cancer was used to select probes for a preferred panel for detection and/or diagnosis of breast cancer. The term "20% or greater positivity" means that if 100 tumor cases were studied, 20 or more of these cases would have shown a presence of the individual marker, while the remaining 80 or fewer cases would not have shown a presence of the individual marker. A preferred panel may include molecular markers selected from Carcinoembrionic Antigen (CEA), C-erb-B2, Cyclin E, E6/E7, Epidermal Growth Factor Receptor (EGFR), Ki-67, p16, p53, Proliferating Cell Nuclear Antigen (PCNA),

Survivin, Telomerase and Vascular Endothelial Growth Factor. A brief description of the library of probes/markers utilized in the present example is provided below.

Carcinoembryonic Antigen (CEA)

[0559] This carcinoembryonic antigen is a highly glycosylated cell surface protein that is overexpressed in a variety of human tumors and has been used as a tumor marker for disease progression in colorectal cancer patients. Previous reports have found elevated serum CEA levels in patients with cervical cancer, although this did not correlate with disease progression (Borras, G., et al., Tumor antigens CA 19.9, CA 125, and CEA in carcinoma of the uterine cervix. Gynecol Oncol, 1995. 57(2): p. 205-11). Sarandakou and associates (Sarandakou, A., et al., Tumour-associated antigens CEA, CA125, SCC and TPS in gynaecological cancer. Eur J Gynaecol Oncol, 1998. 19(1): p. 73-7) demonstrated that serum CEA levels in cervical cancer patients was significantly increased compared to patients with benign gynecological diseases. Another study reported CEA expression increases in cervical intraepithelial neoplasia (CIN III) and carcinoma in situ (CIS) by immunohistochemical staining even though serum CEA levels of these patients were not elevated. The results suggest that CEA immunostaining may be more sensitive than serum CEA to diagnose cervical dysplasia or cancer at early stage (Tendler, A., H.L. Kaufman, and A.S. Kadish, Increased carcinoembryonic antigen expression in cervical intraepithelial neoplasia grade 3 and in cervical squamous cell carcinoma. Hum Pathol, 2000. 31(11): p. 1357-62).

C-erb-B2

[0560] This oncoprotein is a 185 kD membrane-bound glycoprotein. It is a receptor on the cytoplasmic membrane that is homologous to the epidermal growth factor receptor (c-erb-B1). The c-erb-B2 oncogene was independently discovered by several groups and consequently is referred to by various names, including HER2 and neu (Coussens, L., et al., Tyrosine kinase receptor with extensive homology to EGF receptor shares chromosomal location with neu oncogene. Science, 1985. 230(4730): p. 1132-9; Bargmann, C.I., M.C. Hung, and R.A. Weinberg, The neu oncogene encodes an epidermal growth factor receptor-related protein. Nature, 1986. 319(6050): p. 226-30). Over-expression of c-erb-B2 has been demonstrated in 14% to 38% of patients with cervical cancer and has been found to be associated with poor prognosis (Hale, R.J., et al., Prognostic value of c-erbB-2 expression in uterine cervical carcinoma. J Clin Pathol, 1992. 45(7): p. 594-6). Other studies (Sharma, A., et al., Frequent

amplification of C-erbB2 (HER-2/Neu) oncogene in cervical carcinoma as detected by non-fluorescence in situ hybridization technique on paraffin sections. Oncology, 1999. 56(1): p. 83-7; Mitra, A.B., et al., ERBB2 (HER2/neu) oncogene is frequently amplified in squamous cell carcinoma of the uterine cervix. Cancer Res, 1994. 54(3): p. 637-9) found frequent amplification of c-erb-B2 in cervical cancer. C-erb-B2 expression is elevated in cervical carcinoma measured by enzyme-linked immunosorbent assay (ELISA) and immunohistochemistry (IHC) (Kim, J.W., Y.T. Kim, and D.K. Kim, Correlation between EGFR and c-erbB-2 oncoprotein status and response to neoadjuvant chemotherapy in cervical carcinoma. Yonsei Med J, 1999. 40(3): p. 207-14; Ngan, H.Y., et al., Abnormal expression of epidermal growth factor receptor and c-erbB2 in squamous cell carcinoma of the cervix: correlation with human papillomavirus and prognosis. Tumour Biol, 2001. 22(3): p. 176-83).

Cyclin E

[0561] Proteins known as cyclins and an associated group of regulatory proteins called cyclin-dependent kinases (CDKs) regulate the key checkpoints. Cyclin E is a 50 kD protein that complexes with CDK2 in late G1 phase of the cell cycle. Carcinogenesis is characterized by deregulation of the cell cycle. Although p53 is still the most important cell cycle regulator in human malignancies, there is an increased body of evidence indicating that the aberrant expression of cyclins and cyclin-dependent kinase (CDK) inhibitors is considered one of the most important events in malignant transformation of various human cancers. Cho and associates (Cho, N.H., Y.T. Kim, and J.W. Kim, Correlation between G1 cyclins and HPV in the uterine cervix. Int J Gynecol Pathol, 1997. 16(4): p. 339-47) found that cyclin E expression was absent in normal cervical epithelium but was significantly higher in HPV-positive cases. Another study revealed that patients with either invasive cervical cancer or cervical dysplasia have a significantly higher cyclin E index (CEI) than do the control patients (Tae Kim, Y., et al., Expression of cyclin E and p27(KIP1) in cervical carcinoma. Cancer Lett, 2000. 153(1-2): p. 41-50).

E6/E7

[0562] Human papillomavirus (HPV) infection is associated with cervical cancer. E1 and E2 papillomavirus proteins are expressed at the early stage of infection and regulate DNA replication. The E2 protein activates and represses transcription from different HPV promoters. At some stage when viral DNA gets integrated into the cellular genome, the E2 gene is disrupted

or inactivated. This event leads to a depression of the E6 and E7 viral oncogenes. E6 and E7 influence cell proliferation, gene expression, and progression to malignancy (Rosales, R., M. Lopez-Contreras, and R.R. Cortes, Antibodies against human papillomavirus (HPV) type 16 and 18 E2, E6 and E7 proteins in sera: correlation with presence of papillomavirus DNA. J Med Virol, 2001. 65(4): p. 736-44). Studies have shown that in HPV 16 or 18, E6/E7 mRNA was not detected in benign cervical disease. However, 60% of cervical adenocarcinoma in situ (ACIS) and 24% of cervical adenocarcinoma expressed the HPV 16 oncogene. HPV 18 oncogene expression was detected in 27% of ACIS and in 51% of invasive cervical cancer (Riethdorf, S., et al., Analysis of HPV 16 and 18 E6/E7 oncogene expression in cevical and endometrila glandular neoplasias. Cancer Detection and Prevention, 2000. 24(Supplement 1)). Rosales and associates (Rosales, R., M. Lopez-Contreras, and R.R. Cortes, Antibodies against human papillomavirus (HPV) type 16 and 18 E2, E6 and E7 proteins in sera: correlation with presence of papillomavirus DNA. J Med Virol, 2001. 65(4): p. 736-44) studied 172 women with HPV infection and found that antibodies against the E6 and E7 proteins of HPV 16 were found in 52% and 37% of the patients, respectively. Antibodies against the E6 and E7 proteins of HPV 18 were found in 35% and 45% of the patients, respectively. Another study showed that in HPV16 and HPV18, E6/E7 proteins were detected in 48% of cervicovaginal washings and 29% of sera from patients with cervical cancer using enzyme-linked immunosorbent assay (ELISA).

Epidermal Growth Factor Receptor (EGFR)

[0563] Epidermal growth factor receptor (EGFR) is a transmembrane glycoprotein. Binding with its ligands initiates a chain of events that result in DNA synthesis, cell proliferation, and cell differentiation. Activation of the EGFR has been shown to contribute to the growth and spread of many different types of solid tumors. Up-regulation and over-expression of EGFR has been correlated with many processes related to cancer, including uncontrolled cellular proliferation and prevention of apoptosis (Wu, X., et al., Apoptosis induced by an anti-epidermal growth factor receptor monoclonal antibody in a human colorectal carcinoma cell line and its delay by insulin. J Clin Invest, 1995. 95(4): p. 1897-905). Many epithelial tumors express high EGFR, which is associated with advanced disease and poor clinical prognosis, including cervical and gastric cancers, as well as cancers of the colorectum, head and neck (Salomon, D.S., et al., Epidermal growth factor-related peptides and their receptors in human malignancies. Crit Rev Oncol Hematol, 1995. 19(3): p. 183-232). Kim and associates (Kim, J.W., et al., Expression of

epidermal growth factor receptor in carcinoma of the cervix. Gynecol Oncol, 1996. 60(2): p. 283-7) showed that overexpression of EGFR was found in 29 of 40 (72%) invasive cervical cancers and in 5 of 20 (25%) cervical intraepithelial neoplasia (CIN) patients. Over-expression of EGFR appears to be an unfavorable prognostic factor, regardless of the presence of HPV16/18 (Kedzia, W., et al., [Immunohistochemical examination oncogenic c-erb-b2, egf-r proteins and antioncogenic p53 protein in vulvar cancers HPV-16 positive and negative]. Ginekol Pol, 2000. 71(2): p. 63-9).

Ki-67

[0564] This is a nuclear protein expressed in proliferating normal and neoplastic cells. Ki-67 expression occurs during the phase of the cell cycle designated as late G1, S, M and G2. However during the G0 phase, the antigen cannot be detected (Cattoretti, G., et al., Monoclonal antibodies against recombinant parts of the Ki-67 antigen (MIB 1 and MIB 3) detect proliferating cells in microwave-processed formalin-fixed paraffin sections. J Pathol, 1992. 168(4): p. 357-63). Bar and associates (Bar, J.K., et al., Relations between the expression of p53, c-erbB-2, Ki-67 and HPV infection in cervical carcinomas and cervical dysplasias. Anticancer Res, 2001. 21(2A): p. 1001-6) documented that HPV infection, especially accompanied by increase of proliferative activity in dysplasias may define the cell subpopulation predisposed to malignant process development. This is supported by results indicating Ki-67 activity is found in a higher percentage of patients who are HPV-positive than HPV-negative with carcinomas and dysplasias.

p16

[0565] This gene is a cyclin-dependent kinase inhibitor (CDKI) and it may negatively regulate the cell cycle by acting as a tumor suppressor. Cervical dysplasia is induced by persistent infections through high-risk types of human papillomaviruses (HPVs). Outgrowth of dysplastic lesions is triggered by increasing expression of two viral oncogenes, E6 and E7, which both interact with various cell cycle regulating proteins. Among these is the retinoblastoma gene product pRB, which is inactivated by E7. The pRB product inhibits transcription of the cyclin-dependent kinase inhibitor gene p16(INK4a). Increasing expression of viral oncogenes in dysplastic cervical cells might be reflected by increased expression of p16(INK4a). Sano and associates (Sano, T., et al., Immunohistochemical overexpression of p16 protein associated with intact retinoblastoma protein expression in cervical cancer and cervical intraepithelial

neoplasia. Pathol Int, 1998. 48(8): p. 580-5) demonstrated that strong immuno-reactivity for the p16 protein was observed in both nuclei and cytoplasm of all CIN and invasive cancer cases except several low-grade CIN lesions. Studies also showed that overexpression of p16(INK4a) is a specific marker for dysplastic and neoplastic epithelial cells of the cervix and p16, along with Ki-67 and cyclin E, are complimentary surrogate biomarkers for HPV-related cervical neoplasia (Klaes, R., et al., Overexpression of p16(INK4A) as a specific marker for dysplastic and neoplastic epithelial cells of the cervix uteri. Int J Cancer, 2001. 92(2): p. 276-84; Keating, J.T., T. Ince, and C.P. Crum, Surrogate biomarkers of HPV infection in cervical neoplasia screening and diagnosis. Adv Anat Pathol, 2001. 8(2): p. 83-92).

p53

[0566] This is one of the well-known tumor-suppressor genes. The p53 protein is present in minute amounts in normal cells and tissues, but high concentrations occur in a large number of tumors and tumor cell lines. Hence, it can be detected by immunohisto-chemistry (nuclear staining). The increased concentration in tumor cells may be caused by complexing with other proteins, or by mutation of the p53 protein. The gene for p53 is located on chromosome 17p, a frequent site of allele loss in many tumors. Vassallo and associates (Vassallo, J., et al., *High risk HPV and p53 protein expression in cervical intraepithelial neoplasia*. Int J Gynaecol Obstet, 2000. 71(1): p. 45-8) documented that p53 protein overexpression in CIN is associated with high risk HPV infection. By using Western blot analysis and immunohistochemistry, rearrangement of the p53 gene with overexpressed p53 proteins were found in primary cervical cancer (Sahu, G.R., et al., Rearrangement of p53 gene with overexpressed p53 protein in primary cervical cancer. Oncol Rep, 2002. 9(2): p. 433-7).

Proliferating Cell Nuclear Antigen (PCNA)

[0567] This proliferating cell nuclear antigen (PCNA) is a cofactor for DNA polymerase delta. PCNA is expressed in both S phase of the cell cycle and during periods of DNA synthesis associated with DNA repair. PCNA is expressed in proliferating cells in a wide range of normal and malignant tissues. The location of PCNA is nuclear. Kobayashi and others demonstrated there was intimate correlation between the PCNA and mitotic indexes in severe dysplasia and carcinoma in situ (CIS) (Kobayashi, I., et al., The proliferative activity in dysplasia and carcinoma in situ of the uterine cervix analyzed by proliferating cell nuclear antigen immunostaining and silver-binding argyrophilic nucleolar organizer region staining. Hum

Pathol, 1994. 25(2): p. 198-202; Smela, M., M. Chosia, and W. Domagala, Proliferation cell nuclear antigen (PCNA) expression in cervical intraepithelial neoplasia (CIN). An immunohistochemical study. Pol J Pathol, 1996. 47(4): p. 171-4). Another study showed PCNA appears to be a better marker of immunoreactivity for CIN than Ki-67 (Maeda, M.Y., et al., Relevance of the rates of PCNA, Ki-67 and p53 expression according to the epithelial compartment in cervical lesions. Pathologica, 2001. 93(3): p. 189-95).

Survivin

[0568] This is a 142 amino acid protein and is expressed in the G2/M phase of the cell cycle. It is an inhibitor of apoptosis that is selectively overexpressed in common human cancers, but not in normal tissues, and correlates with aggressive disease and unfavorable outcomes (Lehner, R., et al., Immunohistochemical localization of the IAP protein survivin in bladder mucosa and transitional cell carcinoma. Appl Immunohistochem Mol Morphol, 2002. 10(2): p. 134-8). Survivin mRNA is detected in cervical cancer tissue (Saitoh, Y., Y. Yaginuma, and M. Ishikawa, Analysis of Bcl-2, Bax and Survivin genes in uterine cancer. Int J Oncol, 1999. 15(1): p. 137-41). Immunohisto-chemical localization of survivin in benign cervical mucosa, cervical dysplasia, and invasive squamous cell carcinoma showed that nuclear staining was detected in normal mucosa, low-grade dysplasia, and high-grade dysplasia. Staining intensity was greatest in cases with morphologic evidence of HPV infection (Frost, M., et al., Immunohistochemical localization of survivin in benign cervical mucosa, cervical dysplasia, and invasive squamous cell carcinoma. Am J Clin Pathol, 2002. 117(5): p. 738-44).

Telomerase

[0569] This is a ribonucleorotein enzyme that extends and maintains telomeres of eukaryotic chromosomes. Those cells that do not express telomerase have successively shortened telomeres with each cell division, which ultimately leads to chromosomal instability, aging and cell death. It has been hypothesized that infection with high-risk human papillomaviruses (HPVs), in conjunction with other cellular events, plays a critical role in the development of cervical cancer. Activation of the telomerase enzyme complex that synthesizes telomere repeats has been associated with acquisition of immortal phenotype in vitro and is commonly observed in human cancers (Anderson, S., et al., *Telomerase activation in cervical cancer*. Am J Pathol, 1997. 151(1): p. 25-31). Studies have shown that telomerase is exclusively present in cervical carcinomas and a subset of cervical intraepithelial neoplasia grade III lesions, but not in normal

cervical tissues. Activation of telomerase appears to be associated with high-risk-HPV infection, accumulation of inactive p53 proteins and increased cell proliferation in cervical lesions (Snijders, P.J., et al., Telomerase activity exclusively in cervical carcinomas and a subset of cervical intraepithelial neoplasia grade III lesions: strong association with elevated messenger RNA levels of its catalytic subunit and high-risk human papillomavirus DNA. Cancer Res, 1998. 58(17): p. 3812-8; Nair, P., et al., Telomerase, p53 and human papillomavirus infection in the uterine cervix. Acta Oncol, 2000. 39(1): p. 65-70). Another study shows that telomerase activity was detected in 96% of cervical tumor samples and in 69% of pre-malignant cervical scrapings, but not detected in control hysterectomy samples and in cervical scrapings of normal healthy patients. This indicates telomerase is a very sensitive and specific molecular marker for cervical cancer screening (Reddy, V.G., et al., Telomerase-A molecular marker for cervical cancer screening. Int J Gynecol Cancer, 2001. 11(2): p. 100-6).

Vascular Endothelial Growth Factor (VEGF)

[0570] Vascular endothelia growth factor (VEGF) is an important angiogenesis factor and an endothelial cell-specific mitogen. Angiogenesis plays a critical process in the latter stages of carcinogenesis and tumor progression, and is particularly important in the development of distant metastasis. VEGF is known to be one of the most important inducers of angiogenesis and is upregulated in carcinoma of the cervix. Lopez-Ocejo and associates (Lopez-Ocejo, O., et al., Oncogenes and tumor angiogenesis: the HPV-16 E6 oncoprotein activates the vascular endothelial growth factor (VEGF) gene promoter in a p53 independent manner. Oncogene, 2000. 19(40): p. 4611-20) demonstrated that HPV-16 E6-positive cells generally express high levels of the VEGF message and suggest a possibility that the HPV oncoprotein, E6, may contribute to tumor angiogenesis by direct stimulation of the VEGF gene. Another study demonstrated that expression of VEGF is involved in the promotion of angiogenesis in cervical cancer and plays an important role in early cancer invasion (Kodama, J., et al., Vascular endothelial growth factor is implicated in early invasion in cervical cancer. Eur J Cancer, 1999. 35(3): p. 485-9).

VII. Summary of Examples I - VI

[0571] Examples I-VI described above provide preferred probes/markers to be included in panels for detecting and/or diagnosing lung, colorectal, bladder, prostate, breast, and cervical cancer. Figures 8a-c provide a summary of the preferred probes/markers for each cancer type. Figures 8a-c also identify which markers are useful for generic cancer detection utility as well as

those that are more valuable due to their specificity for a particular cancer type. For example, EGFR and Ki-67 are useful for generic cancer detection. Whereas, BL2-10D1, CD44v3, Collagenase, COX-1, HLA-DR, HSP-90, IL-6, IL-10, Lewis X, NMP-22, TGF-β1, TGF-1I, TGF1II and UBC are useful for detection and/or diagnosis of bladder cancer; AE1/AE3, BCA-225, BRCA-1, CA-15:3, Cathespin D, GCDFP-15, HOX-B3, p65, PR and TGK are useful for detection and/or diagnosis of breast cancer, Cyclin E, E6 and E7 are useful for detection and/or diagnosis of cervical cancer, AKT, amphiregulin, β-catenin, Bax, BPG, Cdk/2/cdc2, cFLIP, Cripto-I, Ephrin-B2, Ephrin-B4, Fas-L, HMGI(Y), hMLH1, Lysozyme, Matrilysin, p68, S100A4 and YB-1 are useful for detection and/or diagnosis of colorectal cancer; C-MET, Cyclin A, FGF-2, Glut-1, Glut-3, HERA, MAGE-1, MAGE-3, Mucin 1, Nm23, p120, SP-1, SP-B, Thrombomodulin and TTF-1 are useful for detection and/or diagnosis of lung cancer; and 34βE12, B72.3, FAS, ID-1, Kallidrein 2, Leu 7, P504S, PAP, PIP and PSA are useful for detection and/or diagnosis of prostate cancer.

WHAT IS CLAIMED IS:

1. A panel for detecting a generic disease state or discriminating between specific disease states using cell-based diagnosis, comprising a plurality of probes each of which specifically binds to a marker associated with a generic or specific disease state, wherein the pattern of binding of the component probes of the panel to cells in a cytology specimen is diagnostic of the presence or specific nature of said disease state.

- 2. The panel of claim 1, wherein said generic disease state is selected from the group consisting of cancer and infectious diseases.
- 3. The panel of claim 2, wherein said cancer is selected from the group consisting of epithelial cell-based cancers, solid tumor-based cancers, secretory tumor based cancers, and blood based cancers.
- 4. The panel of claim 2 wherein said infectious disease is selected from the group consisting of cell-based diseases in which the infectious organism is a virus, bacterium, protozoan, parasite, or fungus.
- 5. The panel of claim 1, wherein said panel is optimized by using weighting factors selected from the group consisting of cost, prevalence of a generic disease state in a geographic location, prevalence of a specific disease state in a geographic location, availability of probes and commercial considerations.
- 6. The panel of claim 1, wherein each of said probes comprises a detectable label.
 - 7. The panel of claim 6, wherein said probes comprise antibodies.
- 8. The panel of claim 6, wherein said label is selected from the group consisting of a chromophore, a fluorophore, a dye, a radioisotope and an enzyme.
- 9. The panel of claim 8, wherein said label is a chromophore detected using electromagnetic radiation selected from the group consisting of beta rays, gamma rays, X rays, ultraviolet radiation, visible light, infrared radiation and microwaves.
- 10. The panel of claim 1, wherein said pattern of binding is detected using photonic microscopy.

The panel of claim 10, wherein said photonic microscopy utilizes at least one electromagnetic radiation selected from the group consisting of gamma rays, X rays, beta rays, ultraviolet radiation, visible light, infrared radiation and microwaves.

- 12. The panel of claim 1, wherein said detecting is for sexually transmitted diseases and said discriminating is between chlamydia, trichomonas, gonorrhea, herpes and syphilis.
- 13. A method of forming a panel for detecting a disease state or discriminating between disease states in a patient using cell-based diagnosis, comprising:
- (a) determining the sensitivity and specificity of binding of probes each of which specifically binds to a member of a library of markers associated with a disease state; and
- (b) selecting a limited plurality of said probes whose pattern of binding is diagnostic for the presence or specific nature of said disease state.
 - 14. The method of claim 13, wherein said determining comprises:
- (a) separately contacting a histological or cytological sample from a patient known to be suffering from said disease and a histological or cytological sample from a patient known not to be suffering from said disease with each of said probes;
- (b) measuring the amount of specific binding of each probe with its complementary disease marker at loci where said marker is known to be present in cells of said samples; and
- (c) correlating each said amount with the presence or specific nature of said disease.
- 15. The method of claim 13, wherein said selecting comprises one or more of statistical analytical methods, pattern recognition methods and neural network analysis.
- 16. The method of claim 13, where said selecting comprises the use of weighting factors.
- 17. A method of detecting a disease or discriminating between disease states comprising:
- (a) contacting a cytological sample suspected of containing abnormal cells characteristic of a disease state with a panel according to claim 1; and
- (b) detecting a pattern of binding of said probes that is diagnostic for the presence or specific nature of said disease state.

18. The method of claim 17, wherein said cytological sample is a cellular sample collected from a body fluid, an epithelial cell-based organ system, a fine needle aspiration or a biopsy.

- 19. The method of claim 18, wherein said cytological sample is sputum.
- 20. A panel for detecting a generic disease state or discriminating between specific disease states using cell-based diagnosis, wherein said panel is formed according to the method of claim 13.
- 21. The panel of claim 1, wherein said disease marker is selected from the group consisting of a morphologic biomarker, a genetic biomarker, a cell cycle biomarker, a molecular biomarker and a biochemical biomarker.
- 22. The panel of claim 3, wherein said epithelial cell-based cancer is from the pulmonary, urinary, gastrointestinal or genital tract.
- 23. The panel of claim 3, wherein said solid tumor-based cancer is selected from the group consisting of a sarcoma, breast cancer, pancreatic cancer, liver cancer, kidney cancer, thyroid cancer, and prostate cancer.
- 24. The panel of claim 3, wherein said secretory tumor-based cancer is selected from the group consisting of a sarcoma, breast cancer, pancreatic cancer, liver cancer, kidney cancer, thyroid cancer, and prostate cancer.
- 25. The panel of claim 3, wherein said blood-based cancer is selected from the group consisting of leukemia and lymphoma.
- 26. The method of claim 18, wherein said body fluid is selected from the group consisting of blood, urine, spinal fluid and lymph.
- 27. The method of claim 18, wherein said epithelial cell based organ system is selected from the group consisting of the pulmonary tract, the urinary tract, the genital tract and the gastrointestinal tract.
- 28. The method of claim 18, wherein said final needle aspiration is from solid tissue types in organs and systems.
- 29. The method of claim 18, wherein said biopsy is from solid tissue types in organs and systems.

30. The method of claim 28, wherein said organs and systems are selected from the group consisting of breast, pancreas, liver, kidney, thyroid, bone marrow, muscle, prostate and lung.

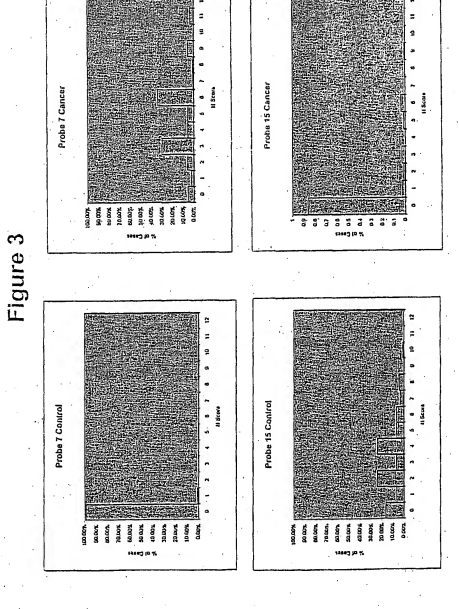
- 31. The panel of claim 21, wherein said morphologic biomarker is selected from the group consisting of DNA ploidy, MACs, and premalignant lesions.
- 32. The panel of claim 21, wherein said genetic biomarker is selected from the group consisting of DNA adducts, DNA mutations and apoptotic indices.
- 33. The panel of claim 21, wherein said cell cycle biomarker is selected from the group consisting of cellular proliferation markers, differentiation markers, regulatory molecules and apoptosis markers.
- 34. The panel of claim 21, wherein said molecular biomarker or biochemical biomarker is selected from the group consisting of oncogenes, tumor suppressor genes, tumor antigens, growth factors and receptors, enzymes, proteins, prostaglandins and adhesion molecules.
- 35. The method of claim 29, wherein said organs and systems are selected from the group consisting of breast, pancreas, liver, kidney, thyroid, bone marrow, muscle, prostate and lung.

Figure 1

3	100.0	94.0	0.18							
Marker	HGF	N-Cadherin	Thrombomodu	lin						
(%)	100.0	100.0	100.0	100.0	97.0	92.0	0.06	85.3	83.0	
Marker	MOC-31	PGF	MUC-1	E-Cadherin	Cyclin A	Bcl-2	Cyclin D1	Rb (loss)	Thyroid	Transcription Factor
(%)	100.0	100.0	100.0	0.96	93.0	87.7	83.5	83.5	80.5	
Marker	VEGF	MUC-1	HGF	EGF Receptor	Telomerase	PCNA	Cyclin A	nın23-H1	Glat1	
(%)	100.0	0.06	86.3	78.3	76.0					
Marker	MOC-31	MUC-1	Telomerase	HGF	Thyroid	Transcription	Factor			
8	100.0	100.0	83.0	82.3	0.08	79.3	79.0	75.5	:	
Marker	Gluti	MOC-31	FGF	Telomerase	PCNA	CD44v6	Cyclin A	HGF		

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Pulmonary Surfactant Pulmonary Surfactant A Thrombomodulin VEGF E-Cadherin Nucleolar Protein N-Cadherin P120 EGF Receptor MAGE-4 Cyclin D1 MAGE-3 Moc-31 FGF CD44v6 Glat3

ligure 2



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Marker	Bladder	Breast	Cervical	Colorectal	Lung	Prostate
mar Kei	Cancer	Cancer	Cancer	Cancer	Cancer	Cancer
EGFR	×	×	X	X	Х	X
Ki-67	X	×	X	Χ .	X	Х
E-Cadherin	. X	X		Х	X	X
p53	X	X	X	X		X
C-erb-B2	×	×	X			· X
Retinoblastoma	×	X		X	X	
Bcl-2		X		X	X	PACINITIAN DATA ASSESSMENT
CEA		X	X	X		
CD44v6	X			, X	X	
Cyclin D1	×			- X	X	
PCNA	×		X	·	Х	
Caveolin-1	X			l X		
-CK 20	X			X		
Cox-2	×			· X		-
ER	• .	X			X	
hMSH2	×		-	X	-	
MDM2	X					X.
MUC-1		X			X.	
N-Cadherin					Х	X
p16		·	X	X		
Survivin	X		X			
Telomerase			X		Х	
VEGF			Х		X	
BL2-10D1	X a					
CD44v3	X					
Collagenase	X		·	-		
Cox-1	X					·
HLA-DR	, X			·		
HSP-90 · ·	X					
IL-6	Х					
IL-10	Х			·		
Lewis X	X					

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Figure 8b

	Bladder	Breast	Cervical	Colorectal	Lung	Prostate
. Marker	Cancer	Cancer	Cancer	Cancer	Cancer	Cancer
NMP-22	X					İ
TGF-β1	X		i			Í
TGF-β1I	X					
TGF-β1II	X		<u> </u>	·		İ
UBC	X		1			
' AE1/AE3		X				
BCA-225		X				
BRCA-1		X				
CA-15.3		X				
Cathepsin D		X				i
GCDFP-15		X		 		
НОХ-ВЗ		X				
p65		X				
PR		Х	j			1
TGK		X				İ
Cyclin E			X		*	
£6			-X		-	
E7			X			·
AKT				X		
Amphiregulin .			1	. X		
β-catenin				X		
Bax				X		
8PG	-			·X		
Cdk2/cdc2				X		
cFLIP				X		
Cripto-l		3-		X		ŀ
Ephrin-82	*			X		
Ephrin-B4				X		1
Fas-L				X		
HMGI(Y)	,			. X-		
hMLH1				X		
Lysozyme				X		
Matrilysin	·			X		
p68				X		
S100A4				×		
Y8-1				X		
C-MET					X	
Cyclin A					X	
FGF-2					Х	
Glut-1					X	

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Figure 8c

Marker	Bladder	Breast	Cervical	Colorectal	Lung	Prostate
	Cancer	Cancer	Cancer	Cancer	Cancer	Cancer
Glut-3					Χ.	
HERA					Х	
MAGE-1		ñ.			Х	·
MAGE-3		±			Х	
Nm23					X	
P120					Χ	
SP-A	*	·	·		Χ	
SP-B					Χ	
Thrombomodulin					X	
TTF-1				·	Χ	
34βE12		-4				Х
B72.3						X
FAS						X
ID-1						X
Kallikrein 2						X ·
Leu 7						X
P504S						X
PAP						Χ
: PIP						X
PSA						X

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